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(54) Title: HEPATITIS C VIRUS REPLICONS AND REPLICON ENHANCED CELLS

(57) Abstract: The present invention features nucleic acid containing one or more adaptive mutations, and HCV replicon enhanced cells. Adaptive mutations are mutations that enhance HCV replicon activity. HCV replicon enhanced cells are cells having an increased ability to maintain an HCV replicon.

# TITLE OF THE INVENTION HEPATITIS C VIRUS REPLICONS AND REPLICON ENHANCED CELLS

# **CROSS-REFERENCE TO RELATED APPLICATIONS**

The present application claims priority to U.S. Serial No. 60/263,479, filed January 23, 2001, hereby incorporated by reference herein.

#### **BACKGROUND OF THE INVENTION**

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The references cited in the present application are not admitted to be prior art to the claimed invention.

It is estimated that about 3% of the world's population are infected with the Hepatitis C virus (HCV). (Wasley, et al., 2000. Semin. Liver Dis. 20, 1-16.) Exposure to HCV results in an overt acute disease in a small percentage of cases, while in most instances the virus establishes a chronic infection causing liver inflammation and slowly progresses into liver failure and cirrhosis. (Iwarson, 1994. FEMS Microbiol. Rev. 14, 201-204.) In addition, epidemiological surveys indicate an important role of HCV in the pathogenesis of hepatocellular carcinoma. (Kew, 1994. FEMS Microbiol. Rev. 14, 211-220, Alter, 1995. Blood 85, 1681-1695.)

The HCV genome consists of a single strand RNA of about 9.5 kb in 20 length, encoding a precursor polyprotein of about 3000 amino acids. (Choo, et al., 1989. Science 244, 362-364, Choo, et al., 1989. Science 244, 359-362, Takamizawa, et al., 1991. J. Virol. 65, 1105-1113.) The HCV polyprotein contains the viral proteins in the order: C-E1-E2-p7-NS2-NS3-NS4A-NS4B-NS5A-NS5B.

Individual viral proteins are produced by proteolysis of the HCV polyprotein. Host cell proteases release the putative structural proteins C, E1, E2, and p7, and create the N-terminus of NS2 at amino acid 810. (Mizushima, et al., 1994. J. Virol. 68, 2731-2734, Hijikata, et al., 1993. P.N.A.S. USA 90, 10773-10777.)

The non-structural proteins NS3, NS4A, NS4B, NS5A and NS5B presumably form the virus replication machinery and are released from the polyprotein. A zinc-dependent protease associated with NS2 and the N-terminus of 30 NS3 is responsible for cleavage between NS2 and NS3. (Grakoui, et al., 1993. J. Virol. 67, 1385-1395, Hijikata, et al., 1993. P.N.A.S. USA 90, 10773-10777.) A distinct serine protease located in the N-terminal domain of NS3 is responsible for proteolytic cleavages at the NS3/NS4A, NS4A/NS4B, NS4B/NS5A and NS5A/NS5B junctions. (Barthenschlager, et al., 1993. J. Virol. 67, 3835-3844, Grakoui, et al.,

1993. Proc. Natl. Acad. Sci. USA 90, 10583-10587, Tomei, et al., 1993. J. Virol. 67, 4017-4026.) NS4A provides a cofactor for NS3 activity. (Failla, et al., J. Virol. 1994. 68, 3753-3760, De Francesco, et al., U.S. Patent No. 5,739,002.) NS5A is a highly phosphorylated protein concurring interferon resistance. (De Francesco, et al., 2000. Semin Liver Dis., 20(1), 69-83, Pawlotsky, 1999. J. Viral Hepat. Suppl. 1, 47-48.) NS5B provides an RNA polymerase. (De Francesco, et al., International Publication Number WO 96/37619, Behrens, et al., 1996. EMBO 15, 12-22, Lohmann, et al., 1998. Virology 249, 108-118.)

Lohmann, et al., Science 285, 110-113, 1999, illustrates the ability of a biscistronic HCV replicon to replicate in a hepatoma cell line. The biscistonic HCV replicon contained a neomycin cistron and an NS2-NS5B or an NS3-NS5B cistron. "NS2-NS5B" refers to a NS2-NS3-NS4A-NS4B-NS5A-NS5B polyprotein. "NS3-NS5B" refers to a NS3-NS4A-NS4B-NS5A-NS5B polyprotein.

Bartenschlager, European Patent Application 1 043 399, published

October 11, 2000 (not admitted to be prior art to the claimed invention), describes a cell culture system for autonomous HCV RNA replication and protein expression.

Replication and protein expression is indicated to occur in sufficiently large amounts for quantitative determination. European Patent Application 1 043 399 indicates that prior cell lines or primary cell cultures infected with HCV do not provide favorable circumstances for detecting HCV replication.

#### SUMMARY OF THE INVENTION

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The present invention features nucleic acid containing one or more adaptive mutations, and HCV replicon enhanced cells. Adaptive mutations are mutations that enhance HCV replicon activity. HCV replicon enhanced cells are cells having an increased ability to maintain an HCV replicon.

An HCV replicon is an RNA molecule able to autonomously replicate in a cultured cell and produce detectable levels of one or more HCV proteins. The basic subunit of an HCV replicon encodes for a HCV NS3-NS5B polyprotein along with a suitable 5' UTR-partial core (PC) region and 3' UTR. The 5' UTR-PC region is made up of a 5'UTR region and about 36 nucleotides of the beginning of the core. Additional regions may be present including those coding for HCV proteins or elements such as the complete core, E1, E2, p7 or NS2; and those coding for other types of proteins or elements such as a encephalomyocarditis virus (EMCV) internal ribosome entry site (IRES), a reporter protein or a selection protein.

The present application identifies different adaptive mutations that enhance HCV replicon activity. Enhancing replicon activity brings about at least one of the following: an increase in replicon maintenance in a cell, an increase in replicon replication, and an increase in replicon protein expression.

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Adaptive mutations are described herein by identifying the location of the adaptive mutation with respect to a reference sequence present in a particular region. Based on the provided reference sequence, the same adaptive mutation can be produced in corresponding locations of equivalent regions having an amino acid sequence different than the reference sequence. Equivalent regions have the same function or encode for a polypeptide having the same function.

Replicon enhanced cells are a preferred host for the insertion and expression of an HCV replicon. Replicon enhanced cells are initially produced by creating a cell containing a HCV replicon and then curing the cell of the replicon. The term "replicon enhanced cell" includes cells cured of HCV replicons and progeny of such cells.

Thus, a first aspect of the present invention describes a nucleic acid molecule comprising at least one of the following regions: an altered NS3 encoding region, an altered NS5A encoding region, and an altered EMCV IRES region. The altered region contains one or more adaptive mutations. Reference to the presence of particular adaptive mutation(s) does not exclude other mutations or adaptive mutations from being present. Adaptive mutations are described with reference to either an encoded amino acid sequence or a nucleic acid sequence.

A nucleic acid molecule can be single-stranded or part of a double strand, and can be RNA or DNA. Depending upon the structure of the nucleic acid molecule, the molecule may be used as a replicon or in the production of a replicon. For example, single-stranded RNA having the proper regions can be a replicon, while double-stranded DNA that includes the complement of a sequence coding for a replicon or replicon intermediate may useful in the production of the replicon or replicon intermediate.

Preferred nucleic acid molecules are those containing region(s) from SEQ. ID. NOs. 1, 2, or 3, or the RNA version thereof, with one or more adaptive mutations. Reference to "the RNA version thereof" indicates a ribose backbone and the presence of uracil instead of thymine.

The presence of a region containing an adaptive mutation indicates that at least one such region is present. In different embodiments, for example, adaptive

mutations described herein are present at least in the NS3 region, in the NS5A region, in the NS5A regions, in the EMCV IRES and NS5A regions, in the EMCV and NS5A regions, and in the ECMV IRES, NS3 and NS5A regions.

Another aspect of the present invention describes an expression vector comprising a nucleotide sequence of an HCV replicon or replicon intermediate coupled to an exogenous promoter. Reference to a nucleotide sequence "coupled to an exogenous promoter" indicates the presence and positioning of an RNA promoter such that it can mediate transcription of the nucleotide sequence and that the promoter is not naturally associated with the nucleotide sequence being transcribed. The expression vector can be used to produce RNA replicons.

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Another aspect of the present invention describes a recombinant human hepatoma cell. Reference to a recombinant cell includes an initially produced cell and progeny thereof.

Another aspect of the present invention describes a method of making a HCV replicon enhanced cell. The method involves the steps of: (a) introducing and maintaining an HCV replicon into a cell and (b) curing the cell of the HCV replicon.

Another aspect of the present invention describes an HCV replicon enhanced cell made by a process comprising the steps of: (a) introducing and maintaining an HCV replicon into a cell and (b) curing the cell of the HCV replicon.

Another aspect of the present invention describes a method of making a HCV replicon enhanced cell comprising an HCV replicon. The method involves (a) introducing and maintaining a first HCV replicon into a cell, (b) curing the cell of the replicon, and (c) introducing and maintaining a second replicon into the cured cell, where the second replicon may be the same or different as the first replicon.

Another aspect of the present invention describes an HCV replicon enhanced cell containing a HCV replicon made by the process involving the step of introducing an HCV replicon into an HCV replicon enhanced cell. The HCV replicon introduced into the HCV replicon enhanced cell may be the same or different than the HCV replicon used to produce the HCV replicon enhanced cell. In a preferred embodiment, the HCV replicon introduced into an HCV replicon enhanced cell is the same replicon as was used to produce the enhanced cell.

Another aspect of the present invention describes a method of measuring the ability of a compound to affect HCV activity using an HCV replicon comprising an adaptive mutation described herein. The method involves providing a compound to a cell comprising the HCV replicon and measuring the ability of the

compound to affect one or more replicon activities as a measure of the effect on HCV activity.

Another aspect of the present invention describes a method of measuring the ability of a compound to affect HCV activity using an HCV replicon enhanced cell that comprises an HCV replicon. The method involves providing a compound to the cell and measuring the ability of the compound to effect one or more replicon activities as a measure of the effect on HCV activity.

Other features and advantages of the present invention are apparent from the additional descriptions provided herein including the different examples.

The provided examples illustrate different components and methodology useful in practicing the present invention. The examples do not limit the claimed invention. Based on the present disclosure the skilled artisan can identify and employ other components and methodology useful for practicing the present invention.

#### 15 BRIEF DESCRIPTION OF THE DRAWINGS

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Figures 1A-1G illustrate the nucleic acid sequence for the pHCVNeo.17 coding strand (SEQ. ID. NO. 3). The different regions of pHCVNeo.17 are provided as follows:

1-341: HCV 5' non-translated region, drives translation of the core-neo fusion protein;

20 342-1181: Core-neo fusion protein, selectable marker;

1190-1800: Internal ribosome entry site of the encephalomyocarditis virus, drives translation of the HCV NS region;

1801-7755: HCV polyprotein from non-structural protein 3 to non-structural protein 5B;

25 1801-3696: Non-structural protein 3 (NS3), HCV NS3 protease/helicase;

3697-3858: Non-structural protein 4A (NS4A), NS3 protease cofactor;

3859-4641: Non-structural protein 4B (NS4B);

4642-5982: Non-structural protein 5A (NS5A);

5983-7755: Non-structural protein 5B (NS5B); RNA-dependent RNA polymerase

30 7759-7989: HCV 3'non-translated region; and

7990-10690 plasmid sequences comprising origin of replication, beta lactamase coding sequence, and T7 promoter.

#### DETAILED DESCRIPTION OF THE INVENTION

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HCV replicons and HCV replicon enhanced cells can be used to produce a cell culture providing detectable levels of HCV RNA and HCV protein. HCV replicons and HCV replicon enhanced hosts can both be obtained by selecting for the ability to maintain an HCV replicon in a cell. As illustrated in the examples provided below, adaptive mutations present in HCV replicons and host cells can both assist replicon maintenance in a cell.

The detectable replication and expression of HCV RNA in a cell culture system has a variety of different uses including being used to study HCV replication and expression, to study HCV and host cell interactions, to produce HCV RNA, to produce HCV proteins, and to provide a system for measuring the ability of a compound to modulate one or more HCV activities.

Preferred cells for use with a HCV replicon are Huh-7 cells and Huh-7 derived cells. "Huh-7 derived cells" are cell produced starting with Huh-7 cells and introducing one or more phenotypic and/or genotypic modifications.

# **Adaptive Mutations**

Adaptive mutations enhance the ability of an HCV replicon to be maintained and expressed in a host cell. Adaptive mutations can be initially selected for using a wild type HCV RNA construct or a mutated HCV replicon. Initial selection involves providing HCV replicons to cells and identifying clones containing a replicon.

Nucleic acid sequences of identified HCV replicons can be determined using standard sequencing techniques. Comparing the sequence of input HCV constructs and selected constructs provides the location of mutations. The effect of particular mutation(s) can be measured by, for example, producing a construct to contain particular mutation(s) and measuring the effect of these mutation(s). Suitable control constructs for comparison purposes include wild type constructs and constructs previously evaluated.

Adaptive mutations were predominantly found in the HCV NS3 and NS5A regions. With the exception of two silent mutations in NS5A and NS5B, consensus mutations occurring in the NS region resulted in changes to the deduced amino acid sequence. Noticeably, the amino acid changes occurred in residues that are conserved in all or a large number of natural HCV isolates. HCV sequences are well known in the art and can be found, for example, in GenBank.

Adaptive mutations described herein can be identified with respect to a reference sequence. The reference sequence provides the location of the adaptive mutation in, for example, the NS3 or NS5A RNA, cDNA, or amino acid sequence. The remainder of the sequence encodes for a functional protein that may have the same, or a different, sequence than the reference sequence.

Preferred NS3 and NS5A adaptive mutations and examples of changes that can be made to produce such mutations are shown in Tables 1 and 2. The amino acid numbering shown in Tables 1 and 2 is with respect to SEQ. ID. NO. 1. The nucleotide numbering shown in Tables 1 and 2 is with respect to SEQ. ID. NO. 2. SEQ. ID. NO. 1 provides the amino acid sequence of the Con1 HCV isolate (Accession Number AJ238799). SEQ. ID. NO. 2 provides the nucleic acid sequence of the Con1 HCV isolate.

TABLE 1

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Preferred NS3 Ad	laptive Mutations
Amino Acid	Nucleotide
· gly1095ala	G3625C
glu1202gly	A3946G
ala1347thr	G4380A

#### TABLE 2

Preferred NS5A	Adaptive Mutations
Amino Acid	Nucleotide
Lys@2039	AAA@6458
asn2041thr	A6463C
ser2173phe	C6859T
ser2197phe	C6931T
leu2198ser	T6934C
ala2199thr	G6936A
ser2204arg	C6953A (or G)

<sup>&</sup>quot;@" refers to an addition.

Preferred adaptive mutations identified with respect to a reference sequence can be produced changing the encoding region of SEQ. ID. NO. 1, or an equivalent sequence, to result in the indicated change. Preferred adaptive mutations provided in Tables 1 and 2 occur in amino acids conserved among different HCV isolates.

Adaptive mutations have different effects. Some mutations alone, or in combination with other mutations, enhance HCV replicon activity. In some cases, two or more mutations led to synergistic effects and in one case, a slightly antagonistic effect was observed.

An adaptive mutation once identified can be introduced into a starting construct using standard genetic techniques. Examples of such techniques are provided by Ausubel, *Current Protocols in Molecular Biology*, John Wiley, 1987-1998, and Sambrook, *et al.*, *Molecular Cloning*, *A Laboratory Manual*, 2<sup>nd</sup> Edition, Cold Spring Harbor Laboratory Press, 1989.

HCV replicons containing adaptive mutations can be built around an NS3 region or NS5A region containing one or more adaptive mutations described herein. The final replicon will contain replicon components needed for replication and may contain additional components.

SEQ. ID. NO. 2 can be used as a reference point for different HCV regions as follows:

5' UTR- nucleotides 1-341;

Core- nucleotides 342-914;

E1- nucleotides 915-1490;

25 E2- nucleotides 1491-2579;

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P7- nucleotides 2580-2768;

NS2- nucleotides 2769-3419;

NS3- nucleotides 3420-5312;

NS4A- nucleotides 5313-5474;

30 NS4B- nucleotides 5475-6257;

NS5A- nucleotides 6258-7598;

NS5B- nucleotides 7599-9371; and

3' UTR- nucleotides 9374-9605.

The amino acid sequences of the different structural and non-structural regions is provided by SEQ. ID. NO. 1.

Nucleic acid sequences encoding for a particular amino acid can be produced taking into account the degeneracy of the genetic code. The degeneracy of the genetic code arises because almost all amino acids are encoded for by different combinations of nucleotide triplets or "codons". The translation of a particular codon into a particular amino acid is well known in the art (see, e.g., Lewin GENES IV, p. 119, Oxford University Press, 1990). Amino acids are encoded for by RNA codons as follows:

A=Ala=Alanine: codons GCA, GCC, GCG, GCU

C=Cys=Cysteine: codons UGC, UGU

10 D=Asp=Aspartic acid: codons GAC, GAU

E=Glu=Glutamic acid: codons GAA, GAG

F=Phe=Phenylalanine: codons UUC, UUU

G=Gly=Glycine: codons GGA, GGC, GGG, GGU

H=His=Histidine: codons CAC, CAU

15 I=Ile=Isoleucine: codons AUA, AUC, AUU

K=Lys=Lysine: codons AAA, AAG

L=Leu=Leucine: codons UUA, UUG, CUA, CUC, CUG, CUU

M=Met=Methionine: codon AUG

N=Asn=Asparagine: codons AAC, AAU

20 P=Pro=Proline: codons CCA, CCC, CCG, CCU

Q=Gln=Glutamine: codons CAA, CAG

R=Arg=Arginine: codons AGA, AGG, CGA, CGC, CGG, CGU

S=Ser=Serine: codons AGC, AGU, UCA, UCC, UCG, UCU

T=Thr=Threonine: codons ACA, ACC, ACG, ACU

25 V=Val=Valine: codons GUA, GUC, GUG, GUU

W=Trp=Tryptophan: codon UGG

Y=Tyr=Tyrosine: codons UAC, UAU.

Constructs, including subgenomic and genomic replicons, containing one or more of the adaptive mutations described herein can also contain additional mutations. The additional mutations may be adaptive mutations and mutations not substantially inhibiting replicon activity. Mutations not substantially inhibiting replicon activity provide for a replicon that can be introduced into a cell and have detectable activity.

#### **HCV Replicon**

HCV replicons include the full length HCV genome and subgenomic constructs. A basic HCV replicon is a subgenomic construct containing an HCV 5' UTR- PC region, an HCV NS3-NS5B polyprotein encoding region, and a HCV 3' UTR. Other nucleic acid regions can be present such as those providing for HCV NS2, structural HCV protein(s) and non-HCV sequences.

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The HCV 5' UTR-PC region provides an internal ribosome entry site (IRES) for protein translation and elements needed for replication. The HCV 5'UTR-PC region includes naturally occurring HCV 5' UTR extending about 36 nucleotides into a HCV core encoding region, and functional derivatives thereof. The 5'-UTR-PC region can be present in different locations such as site downstream from a sequence encoding a selection protein, a reporter, protein, or an HCV polyprotein.

Functional derivatives of the 5'-UTR-PC region able to initiate translation and assist replication can be designed taking into structural requirements for HCV translation initiation. (See, for example, Honda, et al., 1996. Virology 222, 31-42). The affect of different modifications to a 5' UTR-PC region can be determined using techniques that measure replicon activity.

In addition to the HCV 5' UTR-PC region, non-HCV IRES elements can also be present in the replicon. The non-HCV IRES elements can be present in different locations including immediately upstream the region encoding for an HCV polyprotein. Examples of non-HCV IRES elements that can be used are the EMCV IRES, poliovirus IRES, and bovine viral diarrhea virus IRES.

The HCV 3' UTR assists HCV replication. HCV 3' UTR includes naturally occurring HCV 3' UTR and functional derivatives thereof. Naturally occurring 3' UTR's include a poly U tract and an additional region of about 100 nucleotides. (Tanaka, et al., 1996. J. Virol. 70, 3307-3312, Kolykhalov, et al., 1996. J. Virol. 70, 3363-3371.) At least in vivo, the 3' UTR appears to be essential for replication. (Kolykhalov, et al., 2000. J. Virol. 2000 4, 2046-2051.) Examples of naturally occurring 3' UTR derivatives are described by Bartenschlager International Publication Number EP 1 043 399.

The NS3-NS5B polyprotein encoding region provides for a polyprotein that can be processed in a cell into different proteins. Suitable NS3-NS5B polyprotein sequences that may be part of a replicon include those present in different HCV strains and functional equivalents thereof resulting in the processing of NS3-NS5B to

a produce functional replication machinery. Proper processing can be measured for by assaying, for example, NS5B RNA dependent RNA polymerase.

The ability of an NS5B protein to provide RNA polymerase activity can be measured using techniques well known in the art. (See, for example, De Franscesco, et al., International Publication Number WO 96/37619, Behrens, et al., 1996. EMBO 15:12-22, Lohmann, et al., 1998. Virology 249:108-118.) Preferably, the sequence of the active NS5B is substantially similar as that provided in SEQ. ID. NO. 1, or a wild type NS5B such as strains HCV-1, HCV-2, HCV-BK, HCV-J, HCV-N, HCV-H. A substantially similar sequence provides detectable HCV polymerase activity and contains 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, or 15 amino acid alterations to that present in a HCV NS5B polymerase. Preferably, no more than 1, 2, 3, 4 or 5 alterations are present.

Alterations to an amino acid sequence provide for substitution(s), insertion(s), deletion(s) or a combination thereof. Sites of different alterations can be designed taking into account the amino acid sequences of different NS5B polymerases to identify conserved and variable amino acid, and can be empirically determined.

HCV replicons can be produced in a wide variety of different cells and in vitro. Suitable cells allow for the transcription of a nucleic acid encoding for an HCV replicon.

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### **Additional Sequences**

An HCV replicon may contain non-HCV sequences in addition to HCV sequences. The additional sequences should not prevent replication and expression, and preferably serve a useful function. Sequences that can be used to serve a useful function include a selection sequence, a reporter sequence, transcription elements and translation elements.

#### Selection Sequence

A selection sequence in an HCV replicon facilitates the identification of a cell containing the replicon. Selection sequences are typically used in conjunction with some selective pressure that inhibits growth of cells not containing the selection sequence. Examples of selection sequences include sequences encoding for antibiotic resistance and ribozymes.

Antibiotic resistance can be used in conjunction with an antibiotic to select for cells containing replicons. Examples of selection sequences providing for

antibiotic resistance are sequences encoding resistance to neomycin, hygromycin, puromycin, or zeocin.

A ribozyme serving as a selection sequence can be used in conjunction with an inhibitory nucleic acid molecule that prevents cellular growth. The ribozyme recognizes and cleaves the inhibitory nucleic acid.

#### Reporter Sequence

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A reporter sequence can be used to detect replicon replication or protein expression. Preferred reporter proteins are enzymatic proteins whose presence can be detected by measuring product produced by the protein. Examples of reporter proteins include, luciferase, beta-lactamase, secretory alkaline phosphatase, beta-glucuronidase, green fluorescent protein and its derivatives. In addition, a reporter nucleic acid sequence can be used to provide a reference sequence that can be targeted by a complementary nucleic acid. Hybridization of the complementary nucleic acid to its target can be determined using standard techniques.

# **Additional Sequence Configuration**

Additional non-HCV sequences are preferable 5' or 3' of an HCV replicon genome or subgenomic genome region. However, the additional sequences can be located within an HCV genome as long as the sequences do not prevent detectable replicon activity. If desired, additional sequences can be separated from the replicon by using a ribozyme recognition sequence in conjunction with a ribozyme.

Additional sequences can be part of the same cistron as the HCV polyprotein or can be a separate cistron. If part of the same cistron, the selection or reporter sequence coding for a protein should result in a product that is either active as a chimeric protein or is cleaved inside a cell so it is separated from HCV protein.

Selection and reporter sequences encoding for a protein when present as a separate cistron should be associated with elements needed for translation. Such elements include a 5' IRES.

#### **Detection Methods**

Methods for detecting replicon activity include those measuring the production or activity of replicon RNA and encoded for protein. Measuring includes qualitative and quantitative analysis.

Techniques suitable for measuring RNA production include those detecting the presence or activity of RNA. The presence of RNA can be detected using, for example, complementary hybridization probes or quantitative PCR. Techniques for measuring hybridization between complementary nucleic acid and quantitative PCR are well known in the art. (See for example, Ausubel, Current Protocols in Molecular Biology, John Wiley, 1987-1998, Sambrook, et al., Molecular Cloning, A Laboratory Manual, 2<sup>nd</sup> Edition, Cold Spring Harbor Laboratory Press, 1989, and U.S. Patent No. 5,731,148.)

RNA enzymatic activity can be provided to the replicon by using a ribozyme sequence. Ribozyme activity can be measured using techniques detecting the ability of the ribozyme to cleave a target sequence.

Techniques for measuring protein production include those detecting the presence or activity of a produced protein. The presence of a particular protein can be determined by, for example, immunological techniques. Protein activity can be measured based on the activity of an HCV protein or a reporter protein sequence.

Techniques for measuring HCV protein activity vary depending upon the protein that is measured. Techniques for measuring the activity of different nonstructural proteins such as NS2/3, NS3, and NS5B, are well known in the art. (See, for example, references provided in the Background of the Invention.)

Assays measuring replicon activity also include those detecting virion production from a replicon that produces a virion; and those detecting a cytopathic effect from a replicon producing proteins exerting such an effect. Cytopathic effects can be detected by assays suitable to measure cell viability.

Assays measuring replicon activity can be used to evaluate the ability of a compound to modulate HCV activities. Such assays can be carried out by providing one or more test compounds to a cell expressing an HCV replicon and measuring the effect of the compound on replicon activity. If a preparation containing more than one compound is found to modulate replicon activity, individual compounds or smaller groups of compounds can be tested to identify replicon active compounds.

Compounds identified as inhibiting HCV activity can be used to produce replicon enhanced cells and may be therapeutic compounds. The ability of a compound to serve as a therapeutic compound can be confirmed using animal models such as a chimpanzee to measure efficacy and toxicity.

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#### Replicon Enhanced Host Cell

Replicon enhanced cells are initially produced by selecting for a cell able to maintain an HCV replicon and then curing the cell of the replicon. Cells produced in this fashion were found to have an increased ability to maintain a replicon upon subsequent HCV replicon transfection.

Initial transfection can be performed using a wild-type replicon or a replicon containing one or more adaptive mutations. If a wild-type replicon is employed, the replicon should contain a selection sequence to facilitate replicon maintenance.

Cells can be cured of replicons using different techniques such as those employing replicon inhibitory agent. In addition, replication of HCV replicons is substantially reduced in confluent cells. Thus, it is conceivable to cure cells of replicons by culturing them at a high density.

Replicon inhibitory agents inhibit replicon activity or select against a cell containing a replicon. An example of such an agent is IFN-α. Other HCV inhibitory compounds may also be employed. HCV inhibitor compounds are described, for example, in Llinas-Brunet, et al., 2000. Bioorg Med Chem. Lett. 10(20), 2267-2270.

The ability of a cured cell to be a replicon enhanced cell can be measured by introducing a replicon into the cell and determining efficiency of subsequent replicon maintenance and activity.

#### **EXAMPLES**

Examples are provided below to further illustrate different features of the present invention. The examples also illustrate useful methodology for practicing the invention. These examples do not limit the claimed invention.

#### **Example 1: Techniques**

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This example illustrates the techniques employed for producing and analyzing adaptive mutations and replicon enhanced cells.

Manipulation of Nucleic Acids and Construction of Recombinant Plasmids

Manipulation of nucleic acids was done according to standard

protocols. (Sambrook, et al., 1989. Molecular Cloning: A Laboratory Manual, 2<sup>nd</sup> ed.

Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y.) Plasmid DNA was

prepared from ON culture in LB broth using Qiagen 500 columns according to manufacturer instructions.

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Plasmids containing desired mutations were constructed by restriction digestion using restriction sites flanking the mutations or by PCR amplification of the area of interest, using synthetic oligonucleotides with the appropriate sequence. Site directed mutagenesis was carried out by inserting the mutations in the PCR primers. PCR amplification was performed using high fidelity thermostable polymerases or mixtures of polymerases containing a proofreading enzyme. (Barnes, et al., 1994. Proc. Natl. Acad. Sci. 91, 2216-2220.) All plasmids were verified by restriction mapping and sequencing.

pHCVneo17.wt contains the cDNA for an HCV bicistronic replicon identical to replicon I<sub>377</sub>neo/NS3-3'/wt described by Bartenschlager (SEQ. ID. NO. 3) (Lohmann, et al., 1999. Science 285,110-113, EMBL-genbank No. AJ242652). The plasmid comprises the following elements: 5' untranslated region of HCV comprising the HCV-IRES and part of the core (nt1-377); neomycin phosphotransferase coding sequence; and EMCV IRES; HCV coding sequences from NS3 to NS5B; 3' UTR of HCV.

Plasmid pHCVNeo17.GAA is identical to pHCVNeo.17, except that the GAC triplets (nt. 6934-6939 of pHCVNeo17 sequence) coding for the catalytic aspartates of the NS5B polymerase (amino acids 2737 and 2738 of HCV polyprotein) were changed into GCG, coding for alanine.

Plasmid pHCVNeo17.m0 is identical to pHCVNeo17, except that the triplet AGC (nt. 5335-5337 of pHCVNeo17 sequence) coding for the serine of NS5A protein (amino acid 2204 of HCV polyprotein) was changed into AGA, coding for arginine.

Plasmid pHCVNeo17.m1 is identical to pHCVNeo17, except that the triplet AAC (nt. 4846-4848 of pHCVNeo17 sequence) coding for the asparagine of NS5A protein (amino acid 2041 of HCV polyprotein) was changed into ACC, coding for threonine.

Plasmid pHCVNeo17.m2 is identical to pHCVNeo17, except that the triplet TCC (nt. 5242-5244 of pHCVNeo17 sequence) coding for the serine of NS5A protein (amino acid 2173 of HCV polyprotein) was changed into TTC, coding for phenylalanine.

Plasmid pHCVNeo17.m3 is identical to pHCVNeo17, except that the triplet TCC (nt. 5314-5316 of pHCVNeo17 sequence) coding for the serine of NS5A protein (amino acid 2197 of HCV polyprotein) was changed into TTC, coding for phenylalanine.

Plasmid pHCVNeo17.m4 is identical to pHCVNeo17, except that the triplet TTG (nt. 5317-5319 of pHCVNeo17 sequence) coding for the leucine of NS5A protein (amino acid 2198 of HCV polyprotein) was changed into TCG, coding for serine.

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Plasmid pHCVNeo17.m5 is identical to pHCVNeo17, except that an extra triplet AAA coding for lysine was inserted after the triplet GTG (nt. 4840-4843 of pHCVNeo17 sequence), coding for valine 2039 of HCV polyprotein.

Plasmid pHCVNeo17.m6 is identical to pHCVNeo17, except that the triplets GAA and GCC (nt. 2329-2331 and 2764-2766 of pHCVNeo17 sequence) coding for the glutamic acid and the alanine of NS3 protein (amino acid 1202 and 1347 of HCV polyprotein) were changed respectively into GGA and ACC, coding for glycine and threonine. The triplet TCC (nt. 5242-5244 of pHCVNeo17 sequence) coding for the serine of NS5A protein (amino acid 2173 of HCV polyprotein) was changed into TTC, coding for phenylalanine; an extra adenosine was inserted into the EMCV IRES (after the thymidine 1736 of the replicon sequence).

Plasmid pHCVNeo17.m7 is identical to pHCVNeo17, except that the triplet AAC (nt. 4846-4848 of pHCVNeo17 sequence) coding for the asparagine of NS5A protein (amino acid 2041 of HCV polyprotein) was changed into ACC, coding for threonine; the triplet TCC (nt. 5242-5244 of pHCVNeo17 sequence) coding for the serine of NS5A protein (amino acid 2173 of HCV polyprotein) was changed into TTC, coding for phenylalanine.

Plasmid pHCVNeo17.m8 is identical to pHCVNeo17, except that the triplet AAC (nt. 4846-4848 of pHCVNeo17 sequence) coding for the asparagine of NS5A protein (amino acid 2041 of HCV polyprotein) was changed into ACC, coding for threonine; the triplet TCC (nt. 5314-5316 of pHCVNeo17 sequence) coding for the serine of NS5A protein (amino acid 2197 of HCV polyprotein) was changed into TTC, coding for phenylalanine.

Plasmid pHCVNeo17.m9 is identical to pHCVNeo17, except that the triplet AAC (nt. 4846-4848 of pHCVNeo17 sequence) coding for the asparagine of NS5A protein (amino acid 2041 of HCV polyprotein) was changed into ACC, coding

for threonine; the triplet TTG (nt. 5317-5319 of pHCVNeo17 sequence) coding for the leucine of NS5A protein (amino acid 2198 of HCV polyprotein) was changed into TCG, coding for serine.

Plasmid pHCVNeo17.m10 is identical to pHCVNeo17, except that the triplet GAA (nt. 2329-2331 of pHCVNeo17 sequence) coding for the glutamic acid of NS3 protein (amino acid 1202 of HCV polyprotein) was changed into GGA, coding for glycine; an extra triplet AAA coding for lysine was inserted after the triplet GTG (nt. 4840-4843 of pHCVNeo17 sequence), coding for valine 2039 of HCV polyprotein.

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Plasmid pHCVNeo17.m11 is identical to pHCVNeo17, except that the triplet TCC (nt. 5314-5316 of pHCVNeo17 sequence) coding for the serine of NS5A protein (amino acid 2197 of HCV polyprotein) was changed into TTC, coding for phenylalanine. The triplet GCC (nt. 5320-5322 of pHCVNeo17 sequence) coding for the alanine of NS5A protein (amino acid 2199 of HCV polyprotein) was changed into ACC coding for threonine.

Plasmid pHCVNeo17.m12 is identical to pHCVNeo17, except that the triplet AAC (nt. 4846-4848 of pHCVNeo17 sequence) coding for the asparagine of NS5A protein (amino acid 2041 of HCV polyprotein) was changed into ACC, coding for threonine; the triplet TCC (nt. 5314-5316 of pHCVNeo17 sequence) coding for the serine of NS5A protein (amino acid 2197 of HCV polyprotein) was changed into TTC, coding for phenylalanine. The triplet GCC (nt. 5320-5322 of pHCVNeo17 sequence) coding for the alanine of NS5A protein (amino acid 2199 of HCV polyprotein) was changed into ACC coding for threonine.

Plasmid pHCVNeo17.m13 has the same mutations as pHCVNeo17.m8, but also an extra adenosine inserted into the EMCV IRES (after the thymidine 1736 of the replicon sequence).

Plasmid pHCVNeo17.m14 has the same mutations as pHCVNeo17.m11, but also an extra adenosine inserted into the EMCV IRES (after the thymidine 1736 of the replicon sequence).

Plasmid pHCVNeo17.m15 is identical to pHCVNeo17, except that the triplet GCC (nt. 5320-5322 of pHCVNeo17 sequence) coding for the alanine of NS5A protein (amino acid 2199 of HCV polyprotein) was changed into ACC coding for threonine.

Plasmid pRBSEAP.5 is a pHCVNeo.17 derivative where the Neo coding sequence has been replaced with the sequence coding for the human placental alkaline phosphatase corresponding to nucleotides 90-1580 of pBC12/RSV/SEAP plasmid. (Berger, et al., 1988. Gene 66, 1-10.)

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#### RNA Transfection

Transfection was performed using Huh-7 cells. The cells were grown in Dulbecco's modified minimal essential medium (DMEM, Gibco, BRL) supplemented with 10% FCS. For routine work, cells were passed 1 to 5 twice a week using 1x trypsin/EDTA (Gibco, BRL).

Plasmids were digested with the ScaI endonuclease (New England Biolabs) and transcribed *in vitro* with the T7 Megascript kit (Ambion). Transcription mixtures were treated with DNase I (0.1 U/ml) for 30 minutes at 37°C to completely remove template DNA, extracted according to the procedure of Chomczynski (Chomczynski, *et al.*, 1987. *Anal. Biochem. 162*, 156-159), and resuspended with RNase-free phosphate buffered saline (rfPBS, Sambrook, *et al.*, 1989. *Molecular Cloning: A Laboratory Manual*, 2<sup>nd</sup> ed. Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y.).

RNA transfection was performed as described by Liljestrom, et al., 20 1991. J. Virol. 6, 4107-4113, with minor modifications. Subconfluent, actively growing cells were detached from the tissue culture container using trypsin/EDTA. Trypsin was neutralised by addition of 3 to 10 volumes of DMEM/10%FCS and cells were centrifuged for 5 minutes at 1200 rpm in a Haereus table top centrifuge at 40C. Cells were resuspended with ice cold rfPBS by gentle pipetting, counted with a haemocitometer, and centrifuged as above. rfPBS wash was repeated once and cells 25 were resuspended at a concentration of 1-2 x 10<sup>7</sup> cell/ml in rfPBS. Aliquots of cell suspension were mixed with RNA in sterile eppendorf tubes. The RNA/cell mixture was immediately transferred into the electroporation cuvette (precooled on ice) and pulsed twice with a gene pulser apparatus equipped with pulse controller (Biorad). 30 Depending on the experiment, 0.1, 0.2 or 0.4 cm electrode gap cuvettes were used. and settings adjusted (Table 3).

TABLE 3

Cuvette	Volume	Voltage	Capacitance	Resistance	RNA
gap (cm)	(μ1)	(Volts)	(μFa)	(ohm)	(µg)
0.1	70	200	25	infinite	1-10
0.2	200	400	25	infinite	5-20
0.4	800	800	25	infinite	15-100

After the electric shock, cells were left at room temperature for 1-10 5 minutes (essentially the time required to electroporate all samples) and subsequently diluted with at least 20 volumes of DMEM/10%FCS and plated as required for the experiment. Survival and transfection efficiency were monitored by measuring the neutral red uptake of cell cultured for various days in the absence or in the presence of neomycin sulfate (G418). With these parameters, survival of Huh-7 cells was usually 10 40-60% and transfection efficiency ranged between 40% and 100%.

#### Sequence Analysis of Replicon RNAs

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The entire NS region was recloned from 3 different transfection experiments performed with HCVNeo.17 RNA. RNA was extracted from selected clones either using the Qiagen RNAeasy minikit following manufacturer instructions or as described by Chomczynski, et al., 1987. Anal. Biochem. 162, 156-159.

Replicon RNAs (5 µg of total cellular RNA) were retro-transcribed using oligonucleotide HCVG34 (5'- ACATGATCTGCAGAGAGGCCAGT-3'; SEO. ID. No. 4) and the Superscript II reverse transcriptase (Gibco, BRL) according to 20 manufacturer instructions, and subsequently digested with 2 U/ml Ribonuclease H (Gibco BRL). The cDNA regions spanning from the EMCV IRES to the HCV 3' end were amplified by PCR using oligonucleotides HCVG39 (5'-GACASGCTGTGATAWATGTCTCCCCC-3'; SEQ. ID. NO. 5) and CITE3 (5'-TGGCTCTCCAAGCGTATTC -3'; SEQ. ID. NO. 6) and the LA Taq DNA polymerase (Takara LA Taq).

Amplified cDNAs were digested with the KpnI endonuclease (New England Biolabs) and the 5.8 kb fragments were gel purified and ligated to the 5.6 kb vector fragment (purified from plasmid pRBSEAP.5 digested with KpnI) using T4

DNA ligase (New England Biolabs) according to manufacturer instructions. Ligated DNAs were transformed by electroporation in DH10B or JM119 strains of *E. coli*.

In the case of NS5A region, total RNA isolated from 3 clones, (HB77, HB60 and HB68) was extracted and used for RT-PCR.  $5\mu g$  of total RNA plus 20 pmole of AS61 oligo (5'-ACTCTCTGCAGTCAAGCGGCTCA-3', RT antisense oligo; SEQ. ID. NO. 7) were heated 5 minutes at 95°C, then DMSO (5% f.c.), DTT (10 mM f.c.), 1 mM dNTP (1 mM f.c.), 1x Superscript buffer (1 x f.c.), and 10 u Superscript (Gibco) were added to a total volume of 20  $\mu$ l and incubated 3 hours at 42°C.  $2\mu$ l of this RT reaction were used to perform PCR with oligos S39 (5'-

CAGTGGATGAACCGGCTGATA-3', sense; SEQ. ID. NO. 8) or S41 (5'-GGGGCGACGGCATCATGCAAACC-3', sense; SEQ. ID. NO. 9) and B43 (5'-CAGGACCTGCAGTCTGTCAAAGG-3', antisense; SEQ. ID. NO. 10) using Elongase Enzyme Mix (Gibco) according the instruction provided by the manufacturer. The resulting PCR fragment was cloned in pCR2.1 vector using the
 TA Cloning kit (Invitrogen) and transformed in Top10F' bacterial strain.

Plasmid DNA was prepared from ON culture of the resulting ampicillin resistant colonies using Qiagen 500 columns according to manufacturer instructions. The presence of the desired DNA insert was ascertained by restriction digestion, and the nucleotide sequence of each plasmid was determined by automated sequencing. Nucleotide sequences and deduced amino acids sequences were aligned using the GCG software.

#### **TaqMan**

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TaqMan analysis was typically performed using 10 ng of RNA in a

reaction mix (TaqMan Gold RT-PCR kit, Perkin Elmer Biosystems) either with HCV

specific oligos/probe (oligo 1: 5'-CGGGAGAGCCATAGTGG-3'; SEQ. ID. NO. 11,

oligo 2: 5'-AGTACCACAAGGCCTTTCG-3'; SEQ. ID. NO. 12, probe: 5'
CTGCGGAACCGGTGAGTACAC-3'; SEQ. ID. NO. 13) or with human GAPDH

specific oligos/probe (Pre-Developed TaqMan Assay Reagents, Endogenous Control

Human GAPDH, Part Number 4310884E, Perkin Elmer Biosystems). PCR was

performed using a Perkin Elmer ABI PRISM 7700 under the following conditions: 30

minutes at 48°C (the RT step), 10 minutes at 95°C and 40 cycles: 15 seconds at 95°C

and 1 minute at 60°C. Quantitative calculations were obtained using the Comparative

C<sub>T</sub> Method (described in User Bulletin #2, ABI PRISM 7700 Sequence Detection

System, Applied Biosystem, Dec 1997) considering the level of GAPDH mRNA

constant. All calculations of HCV RNA are expressed as fold difference over a specific control.

#### Antibodies and Immunological Techniques

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Mouse monoclonal antibody (anti-NS3 mab10E5/24) were produced by standard techniques. (Galfré and Milstein, 1981. *Methods in Enzymology 73*, 1-46.) Purified recombinant protein was used as an immunogen. (Gallinari, *et al.*, 1999. *Biochemistry 38*, 5620-5632.)

For Cell-ELISA analysis, transfected cells were monitored for expression of the NS3 protein by ELISA with the anti-NS3 mab 10E5/24. Cells were seeded into 96 well plates at densities of 40,000, 30,000, 15,000 and 10,000 cells per well and fixed with ice-cold isopropanol at 1, 2, 3 and 4 days post-transfection, respectively. The cells were washed twice with PBS, blocked with 5% non-fat dry milk in PBS + 0.1% Triton X100 + 0.02% SDS (PBSTS) and then incubated overnight at 4°C with 10E5/24 mab diluted 1:2000 in Milk/PBSTS. After washing 5 times with PBSTS, the cells were incubated for 3 hours at room temperature with anti-mouse IgG Fc specific alkaline phosphatase conjugated secondary antibody (Sigma A-7434), diluted 1:2000 in Milk/PBSTS. After washing again as above, the reaction was developed with *p*-nitrophenyl phosphate disodium substrate (Sigma 104-105) and the absorbance at 405 nm read at intervals.

The results were normalized by staining with sulforhodamine B (SRB Sigma S 1402) to determine cell numbers. The alkaline phosphatase substrate was removed from the wells and the cells washed with PBS. The plates were then incubated with 0.4% SRB in 1% acetic acid for 30 minutes (200  $\mu$ l/well), rinsed 4 times in 1% acetic acid, blotted dry and then 200  $\mu$ l/well of 10mM Tris pH 10.5 added. After mixing, the absorbance at 570 nm was read.

# Neutral Red/ Crystal Violet Staining of Foci

The survival of transfected cells in the absence or presence of G418 was monitored by staining of foci/clones with neutral red in vivo with subsequent crystal violet staining. The medium was removed from the cells and replaced with fresh medium containing 0.0025% neutral red (Sigma N2889) and the cells incubated for 3 hours at 37°C. Cells were washed twice with PBS, fixed in 3.5% formaldehyde for 15 minutes, washed twice again in PBS and then with distilled water and the number of (live) foci counted. The cells could then be re-stained with crystal violet

by incubating with an 0.1% crystal violet (Sigma C0775) solution in 20% methanol for 20 minutes at room temperature, followed by 3 washes in 20% methanol and a wash with distilled water.

### 5 Preparation Of Cells Cured Of Endogenous Replicon

Replicon enhanced cells designated 10IFN and Cl.60/cu were produced using different HCV inhibitory agents. Based on the techniques described herein additional replicon enhanced clones can readily be obtained.

10IFN was obtained by curing a Huh-7 cell of a replicon using human IFN-α2b. Huh-7 cells containing HCV replicons (designated HBI10, HBIII4, HBIII27 and HBIII18) were cultured for 11 days in the presence of 100 U/ml recombinant human IFN-α2b (Intron-A, Schering-Plough), and subsequently for 4 days in the absence of IFN-α2b. At several time points during this period, the clones were analyzed for the presence of HCV proteins and RNA by Western and Northern blotting. After 7 days of incubation with IFN-α2b, HCV proteins could no longer be

detected in any of these clones by Western blotting and similar effects were seen with RNA signals in Northern blots. IFN-02b treated cells were stored in liquid nitrogen until used for transfection experiments.

Cl.60/cu was obtained by curing a Huh-7 cell of a replicon using an HCV inhibitory compound. The presence of HCV RNA was determined using PCR (TaqMan) at 4, 9, 12 and 15 days. From day 9 the amount of HCV RNA was below the limit of detection. To further test the disappearance of the replicon, 4 million cells of cured Clone 60 cells (after the 15 days of treatment) were plated in the presence of 1 mg/ml G-418. No viable cells were observed, confirming that absence of HCV replicons able to confer G-418 resistance.

# Example 2: Selection and Characterization of Cell Clones Containing Functional HCV Replicons

Huh-7 cells (2-8x10<sup>6</sup>) were transfected by electroporation with *in vitro*transcribed replicon RNAs (10-20 μg), plated at a density ranging from 2.5x10<sup>3</sup> to
10x10<sup>3</sup>/cm<sup>2</sup>, and cultured in the presence of 0.8-1 mg/ml G418. The majority of
replicon transfected cells became transiently resistant to G418 and duplicated
normally for 7 to 12 days in the presence of the drug, while cells transfected with
irrelevant RNAs and mock transfected cells did not survive more than 7 days (data not
shown). Transient resistance to G418 was likely due to persistence of the Neo protein

expressed from the transfected RNA, since it was observed also with mutated replicons unable to replicate. Approximately 2 weeks after transfection, transient resistance declined, most cells died and small colonies of cells permanently resistant to the antibiotic became visible in samples transfected with HCVNeo.17 RNA, but not in cells transfected with other replicon RNAs.

In several experiments, the frequency of G418 resistant clones ranged between 10 and 100 clones per 10<sup>6</sup> transfected cells. About 20 G418 resistant colonies were isolated, expanded and molecularly characterized. PCR and RT-PCR analysis of nucleic acids indicated that all clones contained HCV RNA but not HCV DNA, demonstrating that G418 resistance was due to the presence of functional replicons (data not shown). This result was confirmed by Northern blot analysis and metabolic labeling with 3H-uridine, which revealed the presence of both genomic and antigenomic HCV RNAs of the expected size (data not shown). Lastly, western blot, immunoprecipitation and immunofluorescence experiments showed that these clones expressed all HCV non-structural proteins as well as Neo protein (data not shown).

Clones differed in terms of cell morphology and growth rate. Replicon RNA copy number (500-10000 molecules/cell) and viral protein expression also varied between different clones (data not shown). However, the amount of replicon RNA and proteins also varied with passages and with culture conditions and was higher when cells were not allowed to reach confluency, suggesting that replicons replicated more efficiently in dividing cells than in resting cells. Processing of the viral polyprotein occurred with kinetics similar to those observed in transfected cells.

Interestingly, in all tested clones HCV replication was efficiently inhibited by treating the cells with IFN-o2b. The EC50 was between 1 and 10 U/ml, depending on the clone.

# **Example 3: Identification of Adaptive Mutations**

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The low number of G418 resistant clones derived from HCVNeo.17 RNA transfection suggested that replication could require mutation(s) capable of adapting the replicon to the host cell (adaptive mutations) and/or that only a small percentage of Huh-7 cells were competent for HCV replication. To verify the first hypothesis, mutations in replicons RNAs derived from selected cell clones were identified.

RNA sequences for different replicons were determined using standard techniques. Such techniques involved isolating RNA from several independent clones, reverse transcription to produce cDNA, amplifying cDNAs by PCR and cloning into an appropriate vector. The cDNA spanning almost the entire HCV NS region (126 bp at the 3' end of the EMCV IRES and 5650 bp of the HCV NS region (i.e., the entire NS ORF and 298 nucleotides at the 3' end) from 5 clones (HB110, HBIII12, HBIII18, HBIII27, HBIV1) were recloned and sequenced. In addition, the NS5A coding region (nt. 4784-6162) from 3 additional clones (HB 77, HB 68 and HB 60) were recloned and sequenced.

To discriminate mutations present in the replicon RNA from those derived from the cloning procedure, at least 2 isolates derived from independent RT-PCR experiments were sequenced for each cell clone. Comparison of the nucleotide sequences with the parental sequence indicated that each isolate contained several mutations (Tables 4A and 4B).

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TABLE 4A

Cell clone	НВІ	II 12	НВІ	II 18	HB	I 10	HBI	II 27
isolate	4	29	28	61	12	43	13	72
	1674-	1674-	1674-	1674-	1674-	1674-	1674-7460	1674-7460
	7460	7460	7460	7460	7460	7460		ì
EMCV IRES 126 bp	A @ 1736	A @ 1736		C 1752 T				T 1678 C
NS3	G 2009 C	A 2330 G	T 2150 C	T 2015 C	T 1811 A	A 2330 G	G 2009 C	G 2009 C
1895 bp	A 2698 G	C 2505 T	C 2196 A	A 2338 G	A 2330 G	A 2882 G	T 2015 C	C 2052 A
	G 2764 A	G 2764 A	T 3023 A	C 2616 T	T 2666 C	T 3673 C	C 2336 G	G 2644 A
		T 3085 C	T 3134 C	A 2664 G	T 3395 C		A 3130 T	C 2803 A
	T 3273 C		C 3267 T	A 3148 G			A 3401 G	T 2823 A
				T 3286 C	•		A 3518 C	T 3692 C
				C 3615 T				
				C 3657 T				
NS4A	T 3790 C		A 3847 G	T 3827 A	T 3742 C		A 3743 G	A 3797 G
161 bp					•			
NS4B	T 3869 C	C 4283 T	G 4300 A	A 4136 G	T 4290 C	A 4053 G	G 3880 A	C 4547 T
782 bp	A 4107 G	C 4429 T		A 4261-G		A 2496 C	T 4200 C	
	T 4185 C			G 4309 A		T 4316 G	A 4366 G	
	A 4428 G		4	A 4449 G				

TABLE 4A

Cell clone	HBI	II 12	HBI	II 18	HB	I 10	HBI	II 27
isolate	4	29	28	61	12	43	. 13	72
	1674-	1674-	1674-	1674-	1674-	1674-	1674-7460	1674-7460
	7460	7460	7460	7460	7460	7460		
NS5A	A 4847 C	G 4728 A	C 5243 T	C 4729 A	A 4694 T	A 4675 G	A 4855 G	A 4888 G
1340 bp	G 5158 A	A 4845 G	A 5486 G	T 4993 C	AAA @ 4842	A 4761 G	C 5006 T	C 4985 T
	G 5175 C	C 5243 T	C 5596 T	G 5095 A	T 5237 C	AAA @ 4842	T 5318 C	T 5030 A
	C 5243 T	G 5512 T	G 5823 A	T 5334 C		T 5368 C	A 5574 G	T 5090 A
	C 5390 T	A 5521 G		A 5374 T			G 5866 A	T 5318 C
1	A 5719 G	A 5600 G		T 5379 A				A 5328 G
1		A 5740 C		T 5480 C				A 5399 G
				A 5513 G				A 5574 G
				T 5977 C				
NS5B	T 6316 C	A 6406 G	T 6074 C	A 6150 G	A 6911 G	A 5986 G	G 6479 C	G 6156 A
1477 bp	T 6589 C	G 6756 A	A 6541 G	A 6218 G		T 6099 C	C 6870 T	G 7434 A
	T 7370 C	G 6963 T	A 6732 G	T 7352 A		C 6141 T	A 7213 G	T 7444 C
			A 7350 T		٠ .	G 6463 A	T 7448 C	
			A 7359.G			C 6849 T		
						T 6865 C		

Clone name and isolate number are indicated in the first and second row, respectively.

The first and the last nucleotide of the region that was recloned and sequenced are indicated in the third row.

Nucleotide (IUB code) substitutions are indicated with the original nucleotide, its position and mutated nucleotide.

Nucleotide(s) insertions are indicated with the nucleotide(s), the symbol @ and the position of the nucleotide preceding insertion.

Numbering refers to the first nucleotide of the replicon sequence (EMBL-genbank No. AJ242652). The region in which mutations are located and the nucleotide length of each region are indicated in the left most column.

Silent mutations are in italic.

Non sense mutations are underlined.

15 Consensus mutations are bold.

**TABLE 4B** 

Cell clone	НВ	IVI	H	3 77	HE	3 68	HB	60
isolate	85	93	10	14	42	1	13	7
	1674- 7460	1674- 7460 ·	4784- 6162	4465- 6162	4784- 6162	4465- 6162	4784-6162	4784-6162
EMCV IRES 126 bp		A @ 1736						·
NS3 1895 bp	A 3403 G	A 2572 G A 3454 G						

**TABLE 4B** 

Cell clone	НВ	IVI	HE	3 77	HI	8 68	HE	60
isolate	85	93	10	14	42	1	13	7
	1674- 7460	1674- 7460	4784- 6162	4465- 6162	4784- 6162	4465- 6162	4784-6162	4784-6162
NS4A 161 bp								
NS4B 782 bp	A 4084 G	C 3892 T	·					
NS5A 1340 bp	C 5315 T G 5431 T T 5751 C	A 5225 G C 5315 T G 5320 A T 5356 A G 5523 A	G 5060 C C 5337 A	A 5161 G C 5337 A A 5459 G	C 5298 T C 5337 A A 5639 G	T 4972 C	A 5414 G T 5601 G	C 5337 G C 5551 T G 5806 A
NS5B 1477 bp	A 6365 G	T 5888 A T 6855 C A 7135 G T 7171 C		·		,		

See Table 4A legend.

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The frequency of mutations ranged between  $1.7 \times 10^{-3}$  and  $4.5 \times 10^{-3}$  (average  $3 \times 10^{-3}$ ). The majority of mutations were nucleotide substitutions, although insertions of 1 or more nucleotides were also observed (Tables 4A and 4B).

Approximately 85% of the mutations found only in 1 isolate (non-consensus) were randomly distributed in the recloned fragment, and possibly include mis-incorporation during the PCR amplifications. Conversely, the remaining 15% of the mutations were common to 2 or more isolates derived from independent RT-PCR experiments (consensus mutations), and presumably reflected mutations present in the template RNA.

Consensus mutations were found in all isolates and were either common to isolates derived from the same clone (consensus A), or to isolates derived from different clones (consensus B). Analysis of additional isolates derived from the same cell clones indicated that consensus A mutations were not always present in all isolates derived from one clone (data not shown). This observation, together with the

presence of consensus B mutations, suggests that, even within a single cell clone, replicons exist as quasi-species of molecules with different sequences.

At variance with non-consensus mutations, consensus mutations were not randomly distributed but were clustered in the regions coding for the NS5A protein (frequency 1 x 10<sup>-3</sup>) and for the NS3 protein (frequency 0.5 x 10<sup>-3</sup>). Only one consensus mutation was found in the region coding for the NS5B protein (frequency 0.1 x 10<sup>-3</sup> nucleotides) and none in the regions coding for NS4A and NS4B. Interestingly, 1 consensus mutation was observed also in the EMCV IRES.

With the exception of 2 silent mutations found in NS5A and NS5B, consensus mutations occurring in the NS region resulted in changes in the deduced amino acid sequence (Tables 5A and 5B). Noticeably, these amino acid changes occurred in residues that are conserved in all or most natural HCV isolates.

Interestingly, clones HB 77 and HB 60 displayed different nucleotide substitutions (C5337A and C5337G, respectively) resulting in the same amino acidic mutation (S 2204 R).

TABLE 5A

Cell clone	'HBI	11 12	HBI	11 18	НВ	I 10	НВ	III 27
isolate	4	29	28	61	12	43	13	72
NS3	G 1095 A A 1347 T	E 1202 G A 1347 T			E 1202 G	E 1202 G	G 1095 A	G 1095 A
NS4A	\							
.NS4B					·.			
NS5A	N 2041 T S 2173 F	S 2173 F	S 2173 F	E 2263	K @ 2039	к @ 2039	8	L 2198 S R 2283 R
NS5B							•	

See Table 4A legend.

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TABLE 5B

Cell clone	HB	IVI	HB	77	·HI	3 68	Н	B 60
isolate	85	93	10	14	42	1	13	7
NS3		,						
NS4A							,	
NS4B								
NS5A	S 2197 F	N 2041 T S 2197 F A 2199 T	S 2204 R	S 2204 R	S 2204 R	A 2199 T	A 2199 T	S 2204 R
NS5B	N 2710 N	N 2710 N						

See Table 4A legend.

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### 5 Example 4: Functional Characterization of Consensus Mutations

The identification of consensus mutations in recloned replicons indicated that replication proficiency of replicon RNAs contained in selected cell clones depended from the presence of such mutations. To substantiate this hypothesis, the effect of several consensus mutations on replication were analyzed.

Consensus mutations found in the NS5A region were more closely analyzed. Consensus mutations were segregated from the non-consensus ones, and pHCVNeo.17 derivatives containing single or multiple consensus mutations were constructed (Table 6).

TABLE 6

Construct		Consensus mutations		G418 cfu/10 <sup>5</sup> transfected cells
	NS3	NS5A	EMCV IRES	
pHCVNeo17.wt				0-3
pHCVNeo17.GAA				0
pHCVNeo17.m0		S2204R		30-130
pHCVNeo17.m1		N2041T	•	0-3
pHCVNeo17.m2		S2173F	•	15-60
pHCVNeo17.m3	•	S2197F	•	160-500
pHCVNeo17.m4		L2198S		30-50

TABLE 6

Construct		G418 cfu/10 <sup>5</sup> transfected		
	NS3	NS5A	EMCV IRES	cells
pHCVNeo17.m5		<u>K@2039</u>		25-55
pHCVNeo17.m6	E1202G; A1347T	S2173F	Extra A	13-100
pHCVNeo17.m7		N2041T; S2173F		0-1
pHCVNeo17.m8		N2041T; S2197F		360-500
pHCVNeo17.m9		N2041T; L2198S		140-170
pHCVNeo17.m10	E1202G	<u>K@2039</u>		1060
pHCVNeo17.m11		S2197F; A2199T		. 900
pHCVNeo17.m12		N2041T; S2197F; A2199T		>1000
pHCVNeo17.m13		N2041T; S2197F	Extra A	100
pHCVNeo17.m14		S2197F; A2199T	Extra A	>500
pHCVNeo17.m15		A2199T	•	300-600

Huh-7 cells (2x10<sup>6</sup>) were transfected with 10 µg of RNA transcribed from the indicated constructs. Approximately 2x10<sup>5</sup> cells were plated in a 10 cm tissue culture dish and cultured with 1 mg/ml G418 for 20 days.

Colonies surviving selection were stained with crystal violet and counted.

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RNAs transcribed *in vitro* from these constructs were transfected in Huh-7 cells and the affect on replication was estimated by counting neomycin resistant colonies (G418 cfu). As shown in Table 6, all but 1 construct containing single consensus mutations showed a significant increase on G418 cfu efficiency, thus indicating that the corresponding mutations improved replication. Noticeably, 2 mutants containing single mutations in NS5A (m3 and m15) were clearly more effective than all other single mutants. Results of mutants containing 2 or more mutations, indicated the presence of a synergistic effect in some combinations (m8, m9, m11 and possibly m10), but also a slightly antagonistic effect in 1 mutant (m7).

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#### Example 5: Replicon Replication in the Absence of Selection

Replication of HCV replicons in the absence of a G418 selection was detected using quantitative PCR (TaqMan). At 24 hours post-transfection a large amount of replicon RNA was detected in cells transfected with all replicons, including the GAA control replicon containing mutations in the catalytic GDD motif of the NS5B polymerase. This result suggested that analysis at very early time points (up to 48 hour post-transfection) essentially measured the input RNA. Northern blot analysis also indicated that after 24 hours the majority of the transfected RNA was degraded intracellularly (data not shown).

Analysis at later time points showed that the amount of replicon RNA was considerably reduced at 4 days and eventually became undetectable (6/8 days) in cells transfected with replicon HCVNeo17.wt, but was still high in cells transfected with replicons m0, m3 and m15 (Table 7). At day six, that the amount of replicon RNA became undetectable in cells transfected with replicon HCVNeo17.wt, m0, and m2, but was detectable in cells transfected with replicon m3 and m15 (Table 7).

TABLE 7

Name	Hu	H7
	RNA equ.	RNA equ.
	day 4	day 6
Wt	l x	1 x
hcvneo17.m0	3 x	1 x
hcvneo17.m2	1 x	1 x
hcvneo17.m3	5 x	3 x
hcvneo17.m15	6 x	5 x

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Persistence of m0, m3 and m15 replicons RNA was abolished by treatment with interferon-α or with an HCV inhibitory compound (data not shown). Moreover, RNA persistence was not observed with mutated replicons carrying the NS5B GAA mutation besides adaptive mutations (data not shown). Taken together, these results demonstrated that quantitative PCR could be used to monitor replication at early times post-transfection, and can be used to evaluate the replication proficiency of replicon RNAs containing mutations.

Comparison of the results shown in Tables 6 and 7, indicated that there was a good correlation between the amount of replicon RNA detected by TaqMan and the G418 cfu efficiency. Nonetheless, some mutants (m2, m3) showed a pronounced effect on G418 cfu efficiency, and little if any effect on early replication as measured by TaqMan PCR, while other mutants (m0) showed the reverse behavior.

# Example 6: HCV Replicon Enhanced Cells

HCV replicon enhanced cells were produced by introducing an HCV replicon into a host, then curing the host of the replicon. Adaptive mutations (or combinations of them) by themselves increased up to 2 orders of magnitude the G418 cfu efficiency and enhanced early replication comparably. Nonetheless, even with the most effective mutants, only a small percentage of transfected cells (<5 %, data not shown) gave rise to G418 resistant clones containing functional replicons. This observation was attributed, at least in part to a low cloning efficiency of Huh-7 cells (data not shown), and only a fraction of Huh-7 cells being competent for replication.

Several clones were cured of endogenous replicons by treating them for about 2 weeks with IFN- $\alpha$  or with a HCV inhibitory compound. Analysis at the end of the treatment showed that neither viral proteins nor replicon RNA could be detected.

Cured cells (10IFN and Cl.60/cu) were transfected with mutated replicons and replication efficiency was determined by counting neomycin resistant clones (10IFN) or by TaqMan (10IFN and Cl.60/cu). As shown in Table 8, for all tested replicons the G418 cfu efficiency in 10IFN cells was at least 5 fold higher than in parental Huh-7 cells. This increase in G418 cfu efficiency was particularly relevant for a subset of mutants (m3, m5, m8, m9, m15).

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TABLE 8

Construct	Consensus mutations			G418 cfu/10 <sup>3</sup> transfected cells	
	· NS3	NS5A	EMCV IRES		
pHCVNeo17.wt				12 - 56	
pHCVNeo17.GAA			i	0	
pHCVNeo17.m0		S2204R		. 180 - 1000	
pHCVNeo17.ml		N2041T		8 - 13	
pHCVNeo17.m2		S2173F		2000	
pHCVNeo17.m3		S2197F		1600 - 3000	
pHCVNeo17.m4		L2198S		190 - 650	
pHCVNeo17.m5		K@2039		1600 - 3000	
pHCVNeo17.m6	E1202G; A1347T	S2173F	extra A	600 - 2000	
pHCVNeo17.m7		N2041T; S2173F		170 - 800	
pHCVNeo17.m8		N2041T; S2197F		> 4000	
pHCVNeo17.m9		N2041T; L2198S		1400 - 3000	
pHCVNeo17.m10	E1202G	K@2039	į	>4000	
pHCVNeo17.ml1		S2197F; A2199T		>4000	

TABLE 8

Construct	Consensus mutations			G418 cfu/10 <sup>5</sup> transfected cells	
	NS3	NS5A	EMCV IRES		
pHCVNeo17.m12		N2041T; S2197F; A21997		>4000	
pHCVNeo17.m13		N2041T; S2197F	extra A	>4000	
pHCVNeo17.m14		S2197F; A2199T	extra A	>4000	
pHCVNeo17.m15		A2199T		> 4000	

10IFN cells (2x10<sup>6</sup>) were transfected with 10 µg of RNA transcribed from the indicated constructs. Approximately 2x10<sup>5</sup> cells were plated in a 10 cm tissue culture dish and cultured with 1 mg/ml G418 for 20 days.

Colonies surviving selection were stained with crystal violet and counted.

Strikingly, the best mutants yielded a number of G418 resistant clones ranging between 20 and 80% of the cell clones which grew in the absence of G418 (data not shown), thus indicating that the majority of 10IFN cells were competent for replication. This result was confirmed by TaqMan analysis (Table 9), in which the fold increase versus the parental Huh-7 cells was very high. The data indicates that replicons carrying adaptive mutations replicate vigorously in replicon enhanced cells such as 10IFN and Cl.60/cu.

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TABLE 9

Name	10IFN		Cl.60/cu.	
	RNA equ.	RNA equ.	RNA equ.	RNA equ.
	Day 4	day 6	day 4	Day 6
Wt	1 x	1 x	1 x	1 x
hcvneo17.m0	46 x	12 x	78 x	512 x
hcvneo17.m2	2 x	2 x	1 x	2 x
hcvneo17.m3	68 x	49 x	19 x	392 x
hcvneo17.m15	247 x	. 80 x	268 x	5518 x

Expression of viral proteins was determined in replicon enhanced cells
using an ELISA assay designed to detect the NS3 protein in transfected cells plated in
96 wells microtiter plates (Cell-ELISA). As shown in Table 10, 24 hours posttransfection cells transfected with all tested replicons expressed low but detectable
levels of the NS3 protein.

TABLE 10

		NS3 arbitrary units			
. •	24 h p.t.		96 h p.t.		
Name	· -	+ IFN	-	+IFN	
Construct					
Mock	1	1	1	1	
pHCVNeo17.wt	3.7	4.2	1.2	1.3	
pHCVNeo17.GAA	3.1	3.2	1.1	1	
pHCVNeo17.m0	3.4	3.2	9.9	0.8	
pHCVNeo17.m3	5.7	4.6	4.7	1.5	
pHCVNeo17.m8	6.6	5.1	15.1	1.4	
pHCVNeo17.m10	8	5.6	9.2	1.8	
pHCVNeo17.m11	8.4	6.2	13.6	1.8	

101FN cells (2x10<sup>6</sup>) were transfected with 10 µg of RNA transcribed from the indicated constructs. Cells were plated in 96 wells microtiter plates as indicated in Example 1.

The early expression shown in Table 10 is likely due to translation of transfected RNA, since it was comparable in all replicons (including that carrying the GAA mutation) and was not affected by IFN-α. At 4 days post-transfection, NS3 expression persisted or increased in cells transfected with replicons carrying consensus mutations, but could not be detected anymore in cells transfected with wt and GAA replicons. In addition, NS3 expression was almost completely abolished when cells were cultured in the presence of IFN-α.

Taken together, these results indicated that the level of NS3 expression reflected the replication rate. Indeed, NS3 expression level (Table 10) paralleled the RNA level measured by TaqMan (Table 9). The high replication proficiency of 10IFN cells was further confirmed by immunofluorescence experiments which showed that more than 50% of cells transfected with replicons m8 and m11 expressed high level of viral proteins, and that expression was almost completely abolished by IFN-α.

#### **Example 7: Replication of Full Length Constructs**

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This example illustrates the ability of a full length HCV genome containing adaptive mutations described herein to replicate in a replicon enhanced host cell. The full length sequence of the HCV isolate Con-1 (EMBL-Genbank No. AJ238799) (plasmid pHCVRBFL.wt) and 2 derivatives containing either the N2041T

Where indicated (+IFN), IFN-α (100 U/ml) was added to the culture medium 4 hours post-transfection. At the indicated times post-transfection, cells were fixed and analyzed by Cell-ELISA.

and S2173 F mutations (plasmid pHCVRBFL.m8) or the S2197F and A2199T mutations (plasmid pHCVRBFL.m11) were used as starting constructs.

RNAs transcribed from the starting constructs were transfected in 10IFN cells and their replication proficiency was assessed by Cell-ELISA, immunofluorescence and TaqMan. Both constructs containing consensus mutations (pHCVRBFL.m8 and pHCVRBFL.m11) replicated, while no sign of replication was observed with the wt. construct (data not shown).

#### Example 8: Replicons with Reporter Gene

This example illustrates an HCV replicon containing adaptive mutations and a reporter gene. A pHCVNeo17.wt derivative where the Neo coding region was substituted with that coding for human placental secretory alkaline phosphatase (pRBSEAP5.wt) and a derivative also containing the N2041T and S2173F mutations (plasmid pRBSEAP5.m8) were constructed. RNAs transcribed from these plasmids were transfected in 10IFN cells and their replication proficiency was assessed by measuring secretion of alkaline phosphatase. Analysis of the kinetics of secretion suggested that only plasmid pRBSEAP5.m8 was competent for replication (data not shown).

# 20 Example 9: SEQ. ID. Nos. 1 and 2

SEQ. ID. NOs. 1 and 2 are provided as follows:

#### **SEQ. ID. NO. 1**

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MSTNPKPQRKTKRNTNRRPQDVKFPGGGQIVGGVYLLPRRGPRLGVRATRKT SERSQPRGRRQPIPKARQPEGRAWAQPGYPWPLYGNEGLGWAGWLLSPRGS RPSWGPTDPRRRSRNLGKVIDTLTCGFADLMGYIPLVGAPLGGAARALAHGV RVLEDGVNYATGNLPGCSFSIFLLALLSCLTIPASAYEVRNVSGVYHVTNDCS NASIVYEAADMIMHTPGCVPCVRENNSSRCWVALTPTLAARNASVPTTTIRR HVDLLVGAAALCSAMYVGDLCGSVFLVAQLFTFSPRRHETVQDCNCSIYPGH VTGHRMAWDMMMNWSPTAALVVSQLLRIPQAVVDMVAGAHWGVLAGLA YYSMVGNWAKVLIVMLLFAGVDGGTYVTGGTMAKNTLGITSLFSPGSSQKIQ LVNTNGSWHINRTALNCNDSLNTGFLAALFYVHKFNSSGCPERMASCSPIDAF AQGWGPITYNESHSSDQRPYCWHYAPRPCGIVPAAQVCGPVYCFTPSPVVVG TTDRFGVPTYSWGENETDVLLLNNTRPPQGNWFGCTWMNSTGFTKTCGGPP

CNIGGIGNKTLTCPTDCFRKHPEATYTKCGSGPWLTPRCLVHYPYRLWHYPC TVNFTIFKVRMYVGGVEHRLEAACNWTRGERCNLEDRDRSELSPLLLSTTEW QVLPCSFTTLPALSTGLIHLHQNVVDVQYLYGIGSAVVSFAIKWEYVLLLFLLL ADARVCACLWMMLLIAQAEAALENLVVLNAASVAGAHGILSFLVFFCAAWY 5 IKGRLVPGAAYALYGVWPLLLLLLALPPRAYAMDREMAASCGGAVFVGLILL TLSPHYKLFLARLIWWLQYFITRAEAHLQVWIPPLNVRGGRDAVILLTCAIHPE LIFTITKILLAILGPLMVLQAGITKVPYFVRAHGLIRACMLVRKVAGGHYVQM ALMKLAALTGTYVYDHLTPLRDWAHAGLRDLAVAVEPVVFSDMETKVITW GADTAACGDIILGLPVSARRGREIHLGPADSLEGQGWRLLAPITAYSQOTRGL 10 LGCIITSLTGRDRNQVEGEVQVVSTATQSFLATCVNGVCWTVYHGAGSKTLA **GPKGPITQMYTNVDQDLVGWQAPPGARSLTPCTCGSSDLYLVTRHADVIPVR** RRGDSRGSLLSPRPVSYLKGSSGGPLLCPSGHAVGIFRAAVCTRGVAKAVDFV PVESMETTMRSPVFTDNSSPPAVPQTFQVAHLHAPTGSGKSTKVPAAYAAOG YKVLVLNPSVAATLGFGAYMSKAHGIDPNIRTGVRTITTGAPITYSTYGKFLA 15 DGGCSGGAYDIIICDECHSTDSTTILGIGTVLDQAETAGARLVVLATATPPGSV TVPHPNIEEVALSSTGEIPFYGKAIPIETIKGGRHLIFCHSKKKCDELAAKLSGLG LNAVAYYRGLDVSVIPTSGDVIVVATDALMTGFTGDFDSVIDCNTCVTQTVD FSLDPTFTIETTTVPQDAVSRSQRRGRTGRGRMGIYRFVTPGERPSGMFDSSVL CECYDAGCAWYELTPAETSVRLRAYLNTPGLPVCQDHLEFWESVFTGLTHID 20 AHFLSQTKQAGDNFPYLVAYQATVCARAQAPPPSWDQMWKCLIRLKPTLHG PTPLLYRLGAVQNEVTTTHPITKYIMACMSADLEVVTSTWVLVGGVLAALAA YCLTTGSVVIVGRIILSGKPAIIPDREVLYREFDEMEECASHLPYIEQGMQLAEQ FKQKAIGLLQTATKQAEAAAPVVESKWRTLEAFWAKHMWNFISGIQYLAGLS TLPGNPAIASLMAFTASITSPLTTQHTLLFNILGGWVAAQLAPPSAASAFVGAG .25 IAGAAVGSIGLGKVLVDILAGYGAGVAGALVAFKVMSGEMPSTEDLVNLLPA ILSPGALVVGVVCAAILRRHVGPGEGAVQWMNRLIAFASRGNHVSPTHYVPE SDAAARVTQILSSLTTTQLLKRLHQWINEDCSTPCSGSWLRDVWDWICTVLTD FKTWLQSKLLPRLPGVPFFSCQRGYKGVWRGDGIMQTTCPCGAQITGHVKNG SMRIVGPRTCSNTWHGTFPINAYTTGPCTPSPAPNYSRALWRVAAEEYVEVT RVGDFHYVTGMTTDNVKCPCQVPAPEFFTEVDGVRLHRYAPACKPLLREEV 30 TFLVGLNQYLVGSQLPCEPEPDVAVLTSMLTDPSHITAETAKRRLARGSPPSL ASSSASQLSAPSLKATCTTRHDSPDADLIEANLLWRQEMGGNITRVESENKVV ILDSFEPLQAEEDEREVSVPAEILRRSRKFPRAMPIWARPDYNPPLLESWKDPD YVPPVVHGCPLPPAKAPPIPPPRRKRTVVLSESTVSSALAELATKTFGSSESSA 35 VDSGTATASPDQPSDDGDAGSDVESYSSMPPLEGEPGDPDLSDGSWSTVSEE

ASEDVVCCSMSYTWTGALITPCAAEETKLPINALSNSLLRHHNLVYATTSRSA
SLRQKKVTFDRLQVLDDHYRDVLKEMKAKASTVKAKLLSVEEACKLTPPHS
ARSKFGYGAKDVRNLSSKAVNHIRSVWKDLLEDTETPIDTTIMAKNEVFCVQ
PEKGGRKPARLIVFPDLGVRVCEKMALYDVVSTLPQAVMGSSYGFQYSPGQR
VEFLVNAWKAKKCPMGFAYDTRCFDSTVTENDIRVEESIYQCCDLAPEARQA
IRSLTERLYIGGPLTNSKGQNCGYRRCRASGVLTTSCGNTLTCYLKAAAACRA
AKLQDCTMLVCGDDLVVICESAGTQEDEASLRAFTEAMTRYSAPPGDPPKPE
YDLELITSCSSNVSVAHDASGKRVYYLTRDPTTPLARAAWETARHTPVNSWL
GNIIMYAPTLWARMILMTHFFSILLAQEQLEKALDCQIYGACYSIEPLDLPQIIQ
RLHGLSAFSLHSYSPGEINRVASCLRKLGVPPLRVWRHRARSVRARLLSQGGR
AATCGKYLFNWAVRTKLKLTPIPAASQLDLSSWFVAGYSGGDIYHSLSRARP
RWFMWCLLLLSVGVGIYLLPNR

## SEQ. ID. NO. 2:

15 gccagccccgattgggggcgacactccaccatagatcactcccctgtgaggaactactgtcttcacgcagaaagcgtcta gccatggcgttagtatgagtgtcgtgcagcctccaggacccccctcccgggagagccatagtggtctgcggaaccggtgagtacaccggaattgccaggaccgggtcctttcttggatcaacccgctcaatgcctggagatttgggcgtgccccgcgagactgctagccgagtagtgttgggtcgcgaaaggccttgtggtactgcctgatagggtgcttgcgagtgccccgggaggt ctcgtagaccgtgcaccatgagcacgaatcctaaacctcaaagaaaaaccaaacgtaacaccaaccgccgcccacagga 20 cgtcaagttcccgggcggtggtcagatcgtcggtggagtttacctgttgccgcgcaggggccccaggttgggtgtgcgcgc gactaggaagacttccgagcggtcgcaacctcgtggaaggcgacaacctatccccaaggctcgccagcccgagggtagg gcctgggctcagcccgggtacccctggcccctctatggcaatgagggcttggggtgggcaggatggctcctgtcaccccgt ggctctcggcctagttggggccccacggacccccggcgtaggtcgcgcaatttgggtaaggtcatcgataccctcacgtgc ggcttcgccgatctcatggggtacattccgctcgtcggcgcccccctagggggcgctgccaggggccctggcgcatggcgt 25 ccgggttctggaggacggcgtgaactatgcaacagggaatctgcccggttgctccttttctatcttccttttggctttgctgtccttgggtagcgctcactcccacgctcgcggccaggaacgctagcgtccccactacgacgatacgacgccatgtcgatttgctc gttggggcggctgctctctgctccgctatgtacgtgggagatctctgcgggatctgttttcctcgtcgcccagctgttcaccttctc 30 gcctcgccggcacgagacagtacaggactgcaattgctcaatatatcccggccacgtgacaggtcaccgtatggcttggga tatgatgatgaactggtcacctacagcagccctagtggtatcgcagttactccggatcccacaagctgtcgtggatatggtgg cgggggcccattggggagtcctagcgggccttgcctactattccatggtggggaactgggctaaggttctgattgtgatgcta ctctttgccggcgttgacgggggaacctatgtgacaggggggacgatggccaaaaaacaccctcgggattacgtcctcttttcaccegggtcatcccagaaaatccagcttgtaaacaccaacggcagctggcacatcaacaggactgccctgaactgcaat 35 gactccctcaacactgggttccttgctgcgctgttctacgtgcacaagttcaactcatctggatgcccagagcgcatggccag

ctg cag ccccatcg acg cgt tcg ctcag gg gg gg gg cccatcactta caat gag tcacacag acc gg accag agg gccttacccccgtgtaacatcggggggatcggcaataaaaccttgacctgcccacggactgcttccggaagcaccccgaggcca cttacacca agtgtggttcggggccttggttgacaccca gatgcttggtccactacccatacaggctttggcactacccctgcgaggagagcgttgtaacctggaggacagggacagatcagagcttagcccgctgctgctgtctacaacggagtggcaggta ttgccctgttccttcaccaccctaccggctctgtccactggtttgatccatctccatcagaacgtcgtggacgtacaatacctgt10 ctgtgcctgcttgtggatgatgctgctgatagctcaagctgaggccgccctagagaacctggtggtcctcaacgcggcatcc gtggccggggcgcatggcattctctccttcctcgtgttcttctgtgctgcctggtacatcaagggcaggctggtccctggggc ggcatatgccctctacggcgtatggccgctactcctgctcctgctggcgttaccaccacgagcatacgccatggaccggga gatggcagcatcgtgcggaggcgcggttttcgtaggtctgatactcttgaccttgtcaccgcactataagctgttcctcgctag 15 gct catatggtggttacaatattttatcaccagggccgaggcacacttgcaagtgtggatccccccctcaacgttcgggggggeogegatgeogteatectecteacgtgegegatecacceagagetaatetttaccateaccaaaatettgetegecatacte atctcaccccactgcgggactgggcccacgcggggcctacgagaccttgcggtggcagttgagcccgtcgtcttctctgatat 20 gggggaggagatacatctgggaccggcagacagccttgaagggcaggggtggcgactcctcgcgcctattacggccta ctcccaacagacgcgaggcctacttggctgcatcatcactagcctcacaggccgggacaggaaccaggtcgagggggag gtccaagtggtctccaccgcaacacatctttcctggcgacctgcgtcaatggcgtgtgttggactgtctatcatggtgccgg ctcaaagacccttgccggcccaaagggcccaatcacccaaatgtacaccaatgtggaccaggacctcgtcggctggcaag 25 cgcccccggggcgcgttccttgacaccatgcacctgcggcagctcggacctttacttggtcacgaggcatgccgatgtcat tccggtgcgccggcggcgacagcaggggggagcctactctcccccaggcccgtctcctacttgaagggctcttcgggcggtccactgctctgcccctcggggcacgctgtgggcatctttcgggctgccgtgtgcacccgaggggttgcgaaggcggtg gactttgtaccegtcgagtctatggaaaccactatgcggtccccggtcttcacggacaactcgtcccctccggccgtaccgc aga cattc cag g t g c c catct a cac g c c c cat a g g t a g c a g a g a catta cag g t g c c g c t a cat a g g t g c c g c t a cat a g g t g c c g c t a cat a g g t g c c g c t a cat a cat c cat g t a g c cat a cat a cat c cat30 gggtataaggtgcttgtcctgaacccgtccgtcgccgccaccctaggtttcggggcgtatatgtctaaggcacatggtatcgaggtggttgctctggggggcgcctatgacatcataatatgtgatgagtgccactcaactgactcgaccactatcctgggcatcgg cacagtcctggaccaagcggagacggctggagcgcgactcgtcgtgctcgccaccgctacgcctccgggatcggtcacc gtgccacatccaaacatcgaggaggtggctctgtccagcactggagaaatccccttttatggcaaagccatccccatcgaga 35 ccatcaagggggggaggcacctcattttctgccattccaagaagaaatgtgatgagctcgccgcgaagctgtccggcctcg

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Other embodiments are within the following claims. While several embodiments have been shown and described, various modifications may be made without departing from the spirit and scope of the present invention.

### WHAT IS CLAIMED IS:

1. A nucleic acid molecule comprising a region selected from the group consisting of:

a) an altered HCV NS3 encoding region coding for one or more NS3 mutations, wherein at least one of said NS3 mutations, identified by reference to the amino acid sequence numbering of SEQ. ID. NO. 1, is selected from the group consisting of:

amino acid 1095 being Ala,

- amino acid 1202 being Gly, and amino acid 1347 being Thr;
  - b) an altered HCV NS5A encoding region coding for one or more NS5A mutations, wherein at least one of said NS5A mutations, identified by reference to the amino acid sequence numbering of SEQ. ID. NO. 1, is selected from the group consisting of:

amino acid 2041 being Thr,

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a Lys insertion between residue 2039 and 2040.

amino acid 2173 being Phe,

amino acid 2197 being Phe,

amino acid 2198 being Ser,

amino acid 2199 being Thr, and

amino acid 2204 being Arg; and

- c) an altered encephalomyocarditis virus (EMCV) internal ribosome entry site (IRES) region containing one or more EMCV IRES mutations, wherein at least one of said EMCV IRES mutations, identified by reference to the nucleotide number of SEQ. ID. NO. 3, is an insertion at nucleotide 1736 of adenine.
- 2. The nucleic acid molecule of claim 1, wherein said nucleic acid molecule comprises said NS5A encoding region.
- 3. The nucleic acid molecule of claim 2, wherein at least two of said NS5A adaptive mutations are present.

4. The nucleic acid molecule of claim 2, further comprising a region encoding for a HCV NS3 region, wherein said NS3 region may be the same or different than said altered NS3 region.

- 5. The nucleic acid molecule of claim 4, wherein said nucleic acid molecule is an HCV replicon comprising a HCV 5' UTR-PC region, said NS3 encoding region, an HCV NS4A encoding region, an HCV NS4B encoding region, said NS5A encoding region, an HCV NS5B encoding region, and a HCV 3' UTR.
- 10 6. The nucleic acid molecule of claim 5, wherein said HCV replicon further comprises a sequence encoding for a reporter protein.
  - 7. The nucleic acid molecule of claim 5, wherein said HCV replicon further comprises a sequence encoding for a selection protein.

8. The nucleic acid molecule of claim 5, wherein said HCV replicon further comprises a HCV core encoding region, a HCV E1 encoding region, a

HCV E2 encoding region, a HCV p7 encoding region, and a HCV NS2 encoding region.

- 9. A nucleic acid molecule comprising a region selected from the group consisting of:
- a) an altered HCV NS3 encoding region containing one or more NS3 mutations, wherein at least one of said NS3 mutations, identified by reference to the nucleotide numbering of SEQ. ID. NO. 2, is selected from the group consisting of: nucleotide 3625 being cytosine, nucleotide 3946 being guanine, nucleotide 4380 being adenine,
- b) an altered HCV NS5A encoding region containing one or more 30 NS5A mutations, wherein at least one of said NS5A mutations, identified by reference to the nucleotide numbering of SEQ. ID. NO. 2, is selected from the group consisting of:
  - an insertion of 3 adenine residues between nucleotide 6458 and 6459, nucleotide 6463 being cytosine,
- 35 nucleotide 6859 being thymine or uracil,

nucleotide 6931 being thymine or uracil, nucleotide 6934 being cytosine, nucleotide 6936 being adenine, and nucleotide 6953 being adenine or guanine; and

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- c) an altered encephalomyocarditis virus (EMCV) internal ribosome entry site (IRES) region containing one or more EMCV IRES mutations, wherein at least one of said EMCV IRES mutations, identified by reference to the nucleotide number of SEQ. ID. NO. 3, is an insertion at nucleotide 1736 of adenine.
- 10. The nucleic acid molecule of claim 9, wherein said molecule comprises said altered NS5A encoding region, and the nucleotide sequence of said altered NS5A region is provided for by bases 6258-7598 of SEQ. ID. NO. 2, or the RNA version thereof, modified with one or more of said NS5A modifications selected from the group consisting of:
- an insertion of 3 adenine residues between nucleotide 6458 and 6459, nucleotide 6463 being cytosine, nucleotide 6859 being thymine or uracil, nucleotide 6931 being thymine or uracil, nucleotide 6934 being cytosine,
  nucleotide 6936 being adenine, and
- nucleotide 6953 being adenine or guanine.
- The nucleic acid molecule of claim 10, wherein said molecule is an HCV replicon comprising a HCV 5' UTR-PC region, a modified HCV NS3-NS5B region, and a HCV 3' UTR, wherein said modified NS3-NS5B region comprises said altered NS5A region.
  - 12. The nucleic acid molecule of claim 11, wherein said 5' UTR-PC region is the RNA version of bases 1-377 of SEQ. ID. NO. 2 and said 3' UTR is the RNA version of bases 9374-9605 of SEQ. ID. NO. 2.
    - 13. The nucleic acid molecule of claim 10, wherein said molecule is an HCV replicon comprising a HCV 5' UTR-PC region, a modified HCV NS3-NS5B region, and a HCV 3' UTR, wherein
- said 5' UTR-PC region is the RNA version of bases 1-377 of SEQ. ID. NO. 2;

said 3' UTR is the RNA version of bases 9374-9605 of SEQ. ID. NO. 2; and said modified NS3-NS5B region consists of the RNA version of bases 3420-9371 of SEQ. ID. NO. 2 modified with one or more modifications selected from the group consisting of:

- nucleotide 4380 being adenine,
  nucleotide 3625 being cytosine,
  nucleotide 3946 being guanine,
  an insertion of 3 adenine residues between nucleotide 6458 and nucleotide 6459,
  nucleotide 6463 being cytosine,
- nucleotide 6859 being uracil,
  nucleotide 6931 being uracil,
  nucleotide 6934 being cytosine,
  nucleotide 6936 being adenine, and
  nucleotide 6953 being adenine or guanine.

- 14. The nucleic acid molecule of claim 13, wherein said replicon is a genomic replicon that further comprises the RNA version of nucleotides 378-3419 of SEQ. ID. NO. 2.
- 20 15. A nucleic acid molecule comprising the nucleic acid base sequence of bases 1-7989 of SEQ. ID. NO. 3, or the RNA version thereof, consisting of one or more different modifications selected from the group consisting of:
  - a) nucleotides 5335-5337 modified to code for arginine;
  - b) nucleotides 5242-5244 modified to code for phenylalanine;
- 25 c) nucleotides 5314-5316 modified to code for phenylalanine:
  - d) nucleotides 5317-5319 modified to code for serine;
  - e) nucleotides coding for lysine inserted after nucleotide 4843;
  - f) nucleotides 2329-2331 modified to code for glycine, nucleotides 2764-2766 modified to code for threonine, nucleotides 5242-5244 modified to code for
- phenylalanine, and an extra adenosine inserted after nucleotide 1736;
  - g) nucleotides 4846-4848 modified to code for threonine, and nucleotides 5242-5244 modified to modified to code for phenylalanine;
  - h) nucleotides 4846-4848 modified to code for threonine, and nucleotides 5314-5316 modified to code for phenylalanine;

- i) nucleotides 4846-4848 modified to code for threonine, and nucleotides 5317-5319 modified to code for serine;
- j) nucleotides 2329-2331 modified to code for glycine, and nucleotides coding for lysine inserted after nucleotides 4843;
- 5 k) nucleotides 5314-5316 modified to code for phenylalanine and nucleotides 5320-5322 modified to code for threonine;
  - nucleotides 4846-4848 modified to code for threonine, nucleotides 5314-5316 modified to code for phenylalanine, and nucleotides 5320-5322 modified to code for threonine;
- m) nucleotides 4846-4848 modified to code for threonine, nucleotides 5314-5316 modified to code for phenylalanine, and an extra adenosine inserted after nucleotide 1736; and
  - n) nucleotides 5314-5316 modified to code for phenylalanine, nucleotides 5320-5322 modified to code for threonine, and an extra adenosine inserted after nucleotide 1736;
- 15 and
  - o) nucleotides 5320-5322 modified to code for threonine.
  - 16. The nucleic acid of claim 15, wherein said one or more different modifications is selected from the group consisting of:
- 20 a) C5337A;
  - b) C5243T or U;
  - c) C5315T or U;
  - d) T or U5318C;
  - e) AAA inserted after 4843;
- 25 f) A2330G, G2764A, C5243T or U, and adenosine inserted 1736;
  - g) A4847C and C5243T or U;
  - h) A4847C and C5315T or U;
  - i) A4847C and T or U5318C;
  - j) A2330G and AAA inserted after 4843;
- 30 k) C5315T or U and G5320A;
  - l) A4847C, C5315T or U, and G5320A;
  - m) A4847C, C5315T or U, and adenosine inserted 1736;
  - n) C5315T or U, G5320A and adenosine inserted 1736; and
  - o) G5320A.

17. The nucleic acid of claim 16, wherein said nucleic acid is RNA and comprises said nucleic acid base sequence.

- 5 18. The nucleic acid of claim 17, wherein said nucleic acid is RNA and consists of said nucleic acid base sequence.
- 19. An expression vector comprising a nucleotide sequence coding for the nucleic acid molecule of any one of claims 1-18, wherein said nucleotide
   sequence is transcriptionally coupled to an exogenous promoter.
  - 20. A recombinant cell human hepatoma cell, wherein said cell comprises the nucleic acid of any one of claims 5-8 and 11-18.
- 15 21. The recombinant cell of claim 20, wherein said hepatoma cell is an Huh-7 cell.
  - 22. The recombinant cell of claim 20, wherein said cell is derived from a Huh-7 cell.
  - 23. A recombinant cell made by a process comprising the step of introducing into a human hepatoma cell the nucleic acid of any one of claims 5-8 and 11-18.
- 25 24. A method of making an HCV replicon enhanced cell comprising the steps of:

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- a) introducing and maintaining a HCV replicon in a cell; and
- b) curing said cell of said HCV replicon to produce said replicon enhanced cell.
- 25. The method of claim 24, wherein said cell is a human hepatoma cell.
- 26. The method of claim 24, wherein said cell is a Huh-7 cell or is derived from a Huh-7 cell.

	27.	The method of claim 26, further comprising the step of
confirming the	ability	of said replicon enhanced cell to maintain an HCV replicon

- 5 28 A method of making an HCV replicon enhanced cell containing a functional HCV replicon comprising the steps of:
  - a) introducing and maintaining a first HCV replicon in a cell;
  - b) curing said cell of said first replicon to produce a cured cell;
- 10 c) introducing and maintaining a second HCV replicon into said cured cell, wherein said second HCV replicon may be the same or different than said first HCV replicon.
- The method of claim 28, wherein said cell is a human hepatoma cell.
  - 30. The method of claim 29, wherein said human hepatoma cell is a Huh-7 cell.
- 20 31. The method of claim 30, wherein said human hepatoma cell is derived from a Huh-7 cell.
  - 32. An HCV replicon enhanced cell made by the method of any one of claims 24-27.
  - 33. An HCV replicon enhanced cell containing a HCV replicon made by the method of any one of claims 28-31.
- 34. A method of measuring the ability of a compound to affect HCV activity comprising the steps of:
  - a) providing said compound to the HCV replicon enhanced cell of claim 33; and
  - b) measuring the ability of said compound to effect one or more replicon activities as a measure of the effect on HCV activity.

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and

35. The method of claim 34, wherein said compound is a ribozyme.

36. The method of claim 34, wherein said compound in an antisense nucleic acid.

- 37. The method of claim 34, wherein compound is an organic compound.
- 38. The method of claim 34, wherein said step (b) measures HCV protein production.
  - 39. The method of claim 33, wherein said step (b) measures production of RNA transcripts.

GCCAGCCCC GATTGGGGGC GACACTCCAC CATAGATCAC TCCCCTGTGA 1 51 GGAACTACTG TCTTCACGCA GAAAGCGTCT AGCCATGGCG TTAGTATGAG 101 TGTCGTGCAG CCTCCAGGAC CCCCCCTCCC GGGAGAGCCA TAGTGGTCTG 151 CGGAACCGGT GAGTACACCG GAATTGCCAG GACGACCGGG TCCTTTCTTG 201 GATCAACCCG CTCAATGCCT GGAGATTTGG GCGTGCCCCC GCGAGACTGC 251 TAGCCGAGTA GTGTTGGGTC GCGAAAGGCC TTGTGGTACT GCCTGATAGG GTGCTTGCGA GTGCCCCGGG AGGTCTCGTA GACCGTGCAC CATGAGCACG AATCCTAAAC CTCAAAGAAA AACCAAAGGG CGCGCCATGA TTGAACAAGA 401 TGGATTGCAC GCAGGTTCTC CGGCCGCTTG GGTGGAGAGG CTATTCGGCT ATGACTGGGC ACAACAGACA ATCGGCTGCT CTGATGCCGC CGTGTTCCGG 501 CTGTCAGCGC AGGGGCGCCC GGTTCTTTTT GTCAAGACCG ACCTGTCCGG TGCCCTGAAT GAACTGCAGG ACGAGGCAGC GCGGCTATCG TGGCTGGCCA 551 601 CGACGGGCGT TCCTTGCGCA GCTGTGCTCG ACGTTGTCAC TGAAGCGGGA 651 AGGGACTGGC TGCTATTGGG CGAAGTGCCG GGGCAGGATC TCCTGTCATC 701 TCACCTTGCT CCTGCCGAGA AAGTATCCAT CATGGCTGAT GCAATGCGGC 751 GGCTGCATAC GCTTGATCCG GCTACCTGCC CATTCGACCA CCAAGCGAAA 801 CATCGCATCG AGCGAGCACG TACTCGGATG GAAGCCGGTC TTGTCGATCA 851 GGATGATCTG GACGAAGAGC ATCAGGGGCT CGCGCCAGCC GAACTGTTCG CCAGGCTCAA GGCGCGCATG CCCGACGGCG AGGATCTCGT CGTGACCCAT 951 GGCGATGCCT GCTTGCCGAA TATCATGGTG GAAAATGGCC GCTTTTCTGG 1001 ATTCATCGAC TGTGGCCGGC TGGGTGTGGC GGACCGCTAT CAGGACATAG CGTTGGCTAC CCGTGATATT GCTGAAGAGC TTGGCGGCGA ATGGGCTGAC 1101 CGCTTCCTCG TGCTTTACGG TATCGCCGCT CCCGATTCGC AGCGCATCGC 1151 CTTCTATCGC CTTCTTGACG AGTTCTTCTG AGTTTAAACA GACCACAACG GTTTCCCTCT AGCGGGATCA ATTCCGCCCC TCTCCCTCCC CCCCCCTAA 1251 CGTTACTGGC CGAAGCCGCT TGGAATAAGG CCGGTGTGCG TTTGTCTATA 1301 TGTTATTTTC CACCATATTG CCGTCTTTTG GCAATGTGAG GGCCCGGAAA CCTGGCCCTG TCTTCTTGAC GAGCATTCCT AGGGGTCTTT CCCCTCTCGC 1351 1401 CAAAGGAATG CAAGGTCTGT TGAATGTCGT GAAGGAAGCA GTTCCTCTGG AAGCTTCTTG AAGACAAACA ACGTCTGTAG CGACCCTTTG CAGGCAGCGG AACCCCCCAC CTGGCGACAG GTGCCTCTGC GGCCAAAAGC CACGTGTATA 1501

# FIG. 1A

1551	AGATACACCT GCAAAGGCGG CACAACCCCA GTGCCACGTT GTGAGTTGGA
1601	TAGTTGTGGA AAGAGTCAAA TGGCTCTCCT CAAGCGTATT CAACAAGGGG
1651	CTGAAGGATG CCCAGAAGGT ACCCCATTGT ATGGGATCTG ATCTGGGGCC
1701	TCGGTGCACA TGCTTTACAT GTGTTTAGTC GAGGTTAAAA AACGTCTAGG
1751	CCCCCGAAC CACGGGGACG TGGTTTTCCT TTGAAAAACA CGATAATACC
1801	ATGGCGCCTA TTACGGCCTA CTCCCAACAG ACGCGAGGCC TACTTGGCTG
1851	CATCATCACT AGCCTCACAG GCCGGGACAG GAACCAGGTC GAGGGGGAGG
. 1901	TCCAAGTGGT CTCCACCGCA ACACAATCTT TCCTGGCGAC CTGCGTCAAT
1951	GGCGTGTGTT GGACTGTCTA TCATGGTGCC GGCTCAAAGA CCCTTGCCGG
2001	CCCAAAGGC CCAATCACCC AAATGTACAC CAATGTGGAC CAGGACCTCG
2051	TCGGCTGGCA AGCGCCCCCC GGGGCGCGTT CCTTGACACC ATGCACCTGC
2101	GGCAGCTCGG ACCTTTACTT GGTCACGAGG CATGCCGATG TCATTCCGGT
2151	GCGCCGGCGG GGCGACAGCA GGGGGAGCCT ACTCTCCCCC AGGCCCGTCT
2201	CCTACTTGAA GGGCTCTTCG GGCGGTCCAC TGCTCTGCCC CTCGGGGCAC
2251	GCTGTGGGCA TCTTTCGGGC TGCCGTGTGC ACCCGAGGGG TTGCGAAGGC
2301	GGTGGACTTT GTACCCGTCG AGTCTATGGA AACCACTATG CGGTCCCCGG
2351	TCTTCACGGA CAACTCGTCC CCTCCGGCCG TACCGCAGAC ATTCCAGGTG
2401	GCCCATCTAC ACGCCCCTAC TGGTAGCGGC AAGAGCACTA AGGTGCCGGC
2451	TGCGTATGCA GCCCAAGGGT ATAAGGTGCT TGTCCTGAAC CCGTCCGTCG
2501	CCGCCACCCT AGGTTTCGGG GCGTATATGT CTAAGGCACA TGGTATCGAC
2551	CCTAACATCA GAACCGGGT AAGGACCATC ACCACGGGTG CCCCCATCAC
2601	GTACTCCACC TATGGCAAGT TTCTTGCCGA CGGTGGTTGC TCTGGGGGCG
2651	CCTATGACAT CATAATATGT GATGAGTGCC ACTCAACTGA CTCGACCACT
2701	ATCCTGGGCA TCGGCACAGT CCTGGACCAA GCGGAGACGG CTGGAGCGCG
2751	ACTCGTCGTG CTCGCCACCG CTACGCCTCC GGGATCGGTC ACCGTGCCAC
2801	ATCCAAACAT CGAGGAGGTG GCTCTGTCCA GCACTGGAGA AATCCCCTTT
2851	TATGGCAAAG CCATCCCCAT CGAGACCATC AAGGGGGGGA GGCACCTCAT
2901	TTTCTGCCAT TCCAAGAAGA AATGTGATGA GCTCGCCGCG AAGCTGTCCG
2951	GCCTCGGACT CAATGCTGTA GCATATTACC GGGGCCTTGA TGTATCCGTC
3001	ATACCAACTA GCGGAGACGT CATTGTCGTA GCAACGGACG CTCTAATGAC
3051	GGGCTTTACC GGCGATTTCG ACTCAGTGAT CGACTGCAAT ACATGTGTCA

# FIG. 1B

CCCAGACAGT CGACTTCAGC CTGGACCCGA CCTTCACCAT TGAGACGACG ACCGTGCCAC AAGACGCGGT GTCACGCTCG CAGCGGCGAG GCAGGACTGG 3151 3201 TAGGGGCAGG ATGGGCATTT ACAGGTTTGT GACTCCAGGA GAACGGCCCT 3251 CGGGCATGTT CGATTCCTCG GTTCTGTGCG AGTGCTATGA CGCGGGCTGT 3301 GCTTGGTACG AGCTCACGCC CGCCGAGACC TCAGTTAGGT TGCGGGCTTA CCTAAACACA CCAGGGTTGC CCGTCTGCCA GGACCATCTG GAGTTCTGGG AGAGCGTCTT TACAGGCCTC ACCCACATAG ACGCCCATTT CTTGTCCCAG 3401 ACTAAGCAGG CAGGAGACAA CTTCCCCTAC CTGGTAGCAT ACCAGGCTAC GGTGTGCGCC AGGGCTCAGG CTCCACCTCC ATCGTGGGAC CAAATGTGGA AGTGTCTCAT ACGGCTAAAG CCTACGCTGC ACGGGCCAAC GCCCCTGCTG 3551 TATAGGCTGG GAGCCGTTCA AAACGAGGTT ACTACCACAC ACCCCATAAC 3651 CAAATACATC ATGGCATGCA TGTCGGCTGA CCTGGAGGTC GTCACGAGCA 3701 CCTGGGTGCT GGTAGGCGGA GTCCTAGCAG CTCTGGCCGC GTATTGCCTG ACAACAGCA GCGTGGTCAT TGTGGGCAGG ATCATCTTGT CCGGAAAGCC 3801 GGCCATCATT CCCGACAGGG AAGTCCTTTA CCGGGAGTTC GATGAGATGG AAGAGTGCGC CTCACACCTC CCTTACATCG AACAGGGAAT GCAGCTCGCC 3851 3901 GAACAATTCA AACAGAAGGC AATCGGGTTG CTGCAAACAG CCACCAAGCA 3951 AGCGGAGGCT GCTGCTCCCG TGGTGGAATC CAAGTGGCGG ACCCTCGAAG 4001 CCTTCTGGGC GAAGCATATG TGGAATTTCA TCAGCGGGAT ACAATATTTA 4051 GCAGGCTTGT CCACTCTGCC TGGCAACCCC GCGATAGCAT CACTGATGGC 4101 ATTCACAGCC TCTATCACCA GCCCGCTCAC CACCCAACAT ACCCTCCTGT TTAACATCCT GGGGGGATGG GTGGCCGCCC AACTTGCTCC TCCCAGCGCT 4151 4201 GCTTCTGCTT TCGTAGGCGC CGGCATCGCT GGAGCGGCTG TTGGCAGCAT 4251 AGGCCTTGGG AAGGTGCTTG TGGATATTTT GGCAGGTTAT GGAGCAGGGG TGGCAGGCGC GCTCGTGGCC TTTAAGGTCA TGAGCGGCGA GATGCCCTCC 4301 4351 ACCGAGGACC TGGTTAACCT ACTCCCTGCT ATCCTCTCC CTGGCGCCCT 4401 AGTCGTCGGG GTCGTGTGCG CAGCGATACT GCGTCGGCAC GTGGGCCCAG 4451 GGGAGGGGC TGTGCAGTGG ATGAACCGGC TGATAGCGTT CGCTTCGCGG 4501 GGTAACCACG TCTCCCCCAC GCACTATGTG CCTGAGAGCG ACGCTGCAGC 4551 ACGTGTCACT CAGATCCTCT CTAGTCTTAC CATCACTCAG CTGCTGAAGA 4601 GGCTTCACCA GTGGATCAAC GAGGACTGCT CCACGCCATG CTCCGGCTCG

FIG. 1C

TGGCTAAGAG ATGTTTGGGA TTGGATATGC ACGGTGTTGA CTGATTTCAA 4701 GACCTGGCTC CAGTCCAAGC TCCTGCCGCG ATTGCCGGGA GTCCCCTTCT TCTCATGTCA ACGTGGGTAC AAGGGAGTCT GGCGGGGCGA CGGCATCATG 4801 CAAACCACCT GCCCATGTGG AGCACAGATC ACCGGACATG TGAAAAACGG TTCCATGAGG ATCGTGGGGC CTAGGACCTG TAGTAACACG TGGCATGGAA CATTCCCCAT TAACGCGTAC ACCACGGGCC CCTGCACGCC CTCCCCGGCG 4901 CCAAATTATT CTAGGGCGCT GTGGCGGGTG GCTGCTGAGG AGTACGTGGA GGTTACGCGG GTGGGGGATT TCCACTACGT GACGGCCATG ACCACTGACA 5001 ACGTAAAGTG CCCGTGTCAG GTTCCGGCCC CCGAATTCTT CACAGAAGTG 5051 GATGGGGTGC GGTTGCACAG GTACGCTCCA GCGTGCAAAC CCCTCCTACG 5101 5151 GGAGGAGGTC ACATTCCTGG TCGGGCTCAA TCAATACCTG GTTGGGTCAC 5201 AGCTCCCATG CGAGCCCGAA CCGGACGTAG CAGTGCTCAC TTCCATGCTC ACCGACCCCT CCCACATTAC GGCGGAGACG GCTAAGCGTA, GGCTGGCCAG GGGATCTCCC CCCTCCTTGG CCAGCTCATC AGCTAGCCAG CTGTCTGCGC 5301 5351 CTTCCTTGAA GGCAACATGC ACTACCCGTC ATGACTCCCC GGACGCTGAC 5401 CTCATCGAGG CCAACCTCCT GTGGCGGCAG GAGATGGGCG GGAACATCAC 5451 CCGCGTGGAG TCAGAAAATA AGGTAGTAAT TTTGGACTCT TTCGAGCCGC 5501 TCCAAGCGGA GGAGGATGAG AGGGAAGTAT CCGTTCCGGC GGAGATCCTG CGGAGGTCCA GGAAATTCCC TCGAGCGATG CCCATATGGG CACGCCCGGA 5601 TTACAACCCT CCACTGTTAG AGTCCTGGAA GGACCCGGAC TACGTCCCTC 5651 CAGTGGTACA CGGGTGTCCA TTGCCGCCTG CCAAGGCCCC TCCGATACCA 5701 CCTCCACGGA GGAAGAGGAC GGTTGTCCTG TCAGAATCTA CCGTGTCTTC TGCCTTGGCG GAGCTCGCCA CAAAGACCTT CGGCAGCTCC GAATCGTCGG 5801 CCGTCGACAG CGGCACGGCA ACGGCCTCTC CTGACCAGCC CTCCGACGAC 5851 GGCGACGCGG GATCCGACGT TGAGTCGTAC TCCTCCATGC CCCCCTTGA GGGGGAGCCG GGGGATCCCG ATCTCAGCGA CGGGTCTTGG TCTACCGTAA 5951 GCGAGGAGGC TAGTGAGGAC GTCGTCTGCT GCTCGATGTC CTACACATGG 6001 ACAGGCGCCC TGATCACGCC ATGCGCTGCG GAGGAAACCA AGCTGCCCAT CAATGCACTG AGCAACTCTT TGCTCCGTCA CCACAACTTG GTCTATGCTA CAACATCTCG CAGCGCAAGC CTGCGGCAGA AGAAGGTCAC CTTTGACAGA 6101 CTGCAGGTCC TGGACGACCA CTACCGGGAC GTGCTCAAGG AGATGAAGGC

FIG. 1D

6201 GAAGGCGTCC ACAGTTAAGG CTAAACTTCT ATCCGTGGAG GAAGCCTGTA AGCTGACGCC CCCACATTCG GCCAGATCTA AATTTGGCTA TGGGGCAAAG GACGTCCGGA ACCTATCCAG CAAGGCCGTT AACCACATCC GCTCCGTGTG 6351 GAAGGACTTG CTGGAAGACA CTGAGACACC AATTGACACC ACCATCATGG 6401 CAAAAATGA GGTTTTCTGC GTCCAACCAG AGAAGGGGGG CCGCAAGCCA GCTCGCCTTA TCGTATTCCC AGATTTGGGG GTTCGTGTGT GCGAGAAAAT GGCCCTTTAC GATGTGGTCT CCACCCTCCC TCAGGCCGTG ATGGGCTCTT 6551 CATACGGATT CCAATACTCT CCTGGACAGC GGGTCGAGTT CCTGGTGAAT 6601 GCCTGGAAAG CGAAGAAATG CCCTATGGGC TTCGCATATG ACACCCGCTG 6651 TTTTGACTCA ACGGTCACTG AGAATGACAT CCGTGTTGAG GAGTCAATCT 6701 ACCAATGTTG TGACTTGGCC CCCGAAGCCA GACAGGCCAT AAGGTCGCTC ACAGAGCGGC TTTACATCGG GGGCCCCCTG ACTAATTCTA AAGGGCAGAA 6801 CTGCGGCTAT CGCCGGTGCC GCGCGAGCGG TGTACTGACG ACCAGCTGCG 6851 GTAATACCCT CACATGTTAC TTGAAGGCCG CTGCGGCCTG TCGAGCTGCG AAGCTCCAGG ACTGCACGAT GCTCGTATGC GGAGACGACC TTGTCGTTAT 6951 CTGTGAAAGC GCGGGGACCC AAGAGGACGA GGCGAGCCTA CGGGCCTTCA 7001 CGGAGGCTAT GACTAGATAC TCTGCCCCCC CTGGGGACCC GCCCAAACCA 7051 GAATACGACT TGGAGTTGAT AACATCATGC TCCTCCAATG TGTCAGTCGC GCACGATGCA TCTGGCAAAA GGGTGTACTA TCTCACCCGT GACCCCACCA 7101 CCCCCTTGC GCGGGCTGCG TGGGAGACAG CTAGACACAC TCCAGTCAAT TCCTGGCTAG GCAACATCAT CATGTATGCG CCCACCTTGT GGGCAAGGAT 7251 GATCCTGATG ACTCATTTCT TCTCCATCCT TCTAGCTCAG GAACAACTTG 7301 AAAAAGCCCT AGATTGTCAG ATCTACGGGG CCTGTTACTC CATTGAGCCA 7351 CTTGACCTAC CTCAGATCAT TCAACGACTC CATGGCCTTA GCGCATTTTC ACTCCATAGT TACTCTCCAG GTGAGATCAA TAGGGTGGCT TCATGCCTCA 7401 GGAAACTTGG GGTACCGCCC TTGCGAGTCT GGAGACATCG GGCCAGAAGT 7451 GTCCGCGCTA GGCTACTGTC CCAGGGGGGG AGGGCTGCCA CTTGTGGCAA 7551 GTACCTCTTC AACTGGGCAG TAAGGACCAA GCTCAAACTC ACTCCAATCC 7601 CGGCTGCGTC CCAGTTGGAT TTATCCAGCT GGTTCGTTGC TGGTTACAGC GGGGGAGACA TATATCACAG CCTGTCTCGT GCCCGACCCC GCTGGTTCAT GTGGTGCCTA CTCCTACTTT CTGTAGGGGT AGGCATCTAT CTACTCCCCA

FIG. 1E

7751 ACCGATGAAC GGGGAGCTAA ACACTCCAGG CCAATAGGCC ATCCTGTTTT THICCCIPIT THITTICIT THITTIPIT THITTIPITITI THITTIPIT 7851 TTCTCCTTTT TTTTTCCTCT TTTTTTCCTT TTCTTTCCTT TGGTGGCTCC ATCTTAGCCC TAGTCACGGC TAGCTGTGAA AGGTCCGTGA GCCGCTTGAC 7951 TGCAGAGAGT GCTGATACTG GCCTCTCTGC AGATCAAGTA CTTCTAGAGA 8001 ATTCTAGCTT GGCGTAATCA TGGTCATAGC TGTTTCCTGT GTGAAATTGT TATCAGCTCA CAATTCCACA CAACATACGA GCCGGAAGCA TAAAGTGTAA 8051 8101 AGCCTGGGAT GCCTAATGAG TGAGCTAACT CACATTAGTT GCGTTGCGCT CACTGCCCGC TTTCCAGTCG GGAAACCTGT CGTGCCAGCT CCATTAGTGA ATCGTCCAAC GCACGGGGAG AGGCGGTTTG CGTATTGGGC GCACTTCCGC TICCTCGCTC ACTGACTCGC TGCGCTCGTT CGTTCGGCTG CGGCGAGCCG TATCAGCTCA CTCAAAGGCG GTAATACGGT TATCCACAGA ATCAGGGGAT AACGCAGGAA AGACCATGTG AGCAAAAGGC CAGCAAAAGG CCAGGAACCG TAAAAAGGCC GCGTTGCTGG CGTTTTTCCA TAGGCTCCGC CCCCTGACG AGCATCACAA AAATCGACGC TCAAGTCAGA GGTGGCGAAA CCCGACAGGA 8501 CTATAAAGAT ACCAGGCGTT TCCCCCTGGA AGCTCCCTCG TGCGCTCTCC 8551 TGTTCCGACC CTGCCGCTTA CCGGATACCT GTCCGCCTTT CTCCCTTCGG 8601 GAAGCGTGGC GCTTTCTCAT AGCTCACGCT GTAGGTATCT CAGTTCGGTG 8651 TAGGTCGTTC GCTCCAAGCT GGGCTGTGTG CACGAACCCC CCGTTCAGCC CGACCGCTGC GCCTTATCCG GTAACTATCG TCTTGAGTCC AACCCGGTAA GACACGACTT ATCGCCACTG GCAGCAGCCA CTGGTAACAG GATTAGCAGA 8801 GCGAGGTATG TAGGCGGTGC TACAGAGTTC TTGAAGTGGT GGCCTAACTA 8851 CGGCTACACT AGAAGGACAG TATTTGGTAT CTGCGCTCTG CTGAAGCCAG TTACCTTCGG AAAAAGAGTT GGTAGCTCTT GATCCGGCAA ACAAACCACC 8951 GCTGGTAGCG GTGGTTTTTT TGTTTGCAAG CAGCAGATTA CGCGCAGAAA 9001 AAAAGGATCT CAAGAAGATC CTTTGATCTT TTCTACGGGG TCTGACGCTC AGTGGAACGA AAACTCACGT TAAGGGATTT TGGTCATGAG ATTATCAAAA 9101 AGGATCTTCA CCTAGATCCT TTTAAATTAA AAATGAAGTT TTAAATCAAT CTAAAGTATA TATGAGTAAA CTTGGTCTGA CAGTTACCAA TGCTTAATCA GTGAGGCACC TATCTCAGCG ATCTGTCTAT TTCGTTCATC CATAGTTGCC 9201 9251 TGACTCCCG TCGTGTAGAT AACTACGATA CGGGAGGGCT TACCATCTGG

FIG. 1F

9301	CCCCAGTGCT GCAATGATAC CGCGAGAACC ACGCTCACCC GCACCAGATT
9351	TATCAGCAAT AAACCAGCCA GCCGGAAGTG CGCTGCGGAG AAGTGGTCCT
9401	GCAACTTTAT CCGCCTCCAT CCAGTCTATT AGTTGTTGCC GGGAAGCTAG
9451	AGTAAGTAGT TCGCCAGTCA GCAGTTTGCG TAACGTCGTT GCCATAGCAA
9501	CAGGCATCGT GGTGTCACGC TCGTCGTTTG GTATGGCTTC ATTCAGCTCC
9551	GGCTCCCAAC GATCAAGGCG AGTTACATGA TCCCCCATGT TGTGCAAAAA
9601	AGCGGTTAGC TCCTTCGGTC CTCCGATCGT TGTCAGAAGT AAGTTGGCCG
9651	CAGTGTTATC ACTCATGGTT ATGGCAGCAC TGCATAATTC TCTTACTGTC
9701	ATGCCATCCG TAAGATGCTT TTCTGTGACT GGTGAGTACT CAACCAAGTC
9751	ATTCTGAGAA TAGTGTATGC GGCGACCGAG TTGCTCTTGC CCGGCGTCAA
9801	TACGGGATAA TACCGCGCCA CATAGCAGAA CTTTAAAAGT GCTCATCATT
9851	GGAAAACGTT CTTCGGGGCG AAAACTCTCA AGGATCTTAC CGCTGTTGAG
9901	ATCCAGTTCG ATGTAACCCA CTCGTGCACC CAACTGATCT TCAGCATCTT
9951	TTACTTTCAC CAGCGTTTCT GGGTGAGCAA AAACAGGAAG GCAAAATGCC
10001	GCAAAAAAGG GAATAAGGGC GACACGGAAA TGTTGAATAC TCATACTCTT
10051	CCTTTTTCAA TATTATTGAA GCATTTATCA GGGTTATTGT CTCATGAGCG
10101	GATACATATT TGAATGTATT TAGAAAAATA AACAAATAGG GGTTCCGCGC
10151	ACATTTCCCC GAAAAGTGCC ACCTGACGTC TAAGAAACCA TTATTACCAT
10201	GACATTAACC TATAAAAATA GGCGTATCAC GAAGCCCTTT CGTCTAGCGC
10251	GTTTCGGTGA TGACGGTGAA AACCTCTGAC ACTTGCAGCT CCCGCAGACG
10301	GTCACAGCTT GTCTGTAAGC GGATGCCGGG AGCAGGCAAG CCCGTCAGGG
10351	CGCGTCAGTG GGTGTTGGCG GGTGTCGGGG CTGGCTTAAC TATGCGGCAT
10401	CAGAGCAGAT TGTACTGAGA GTACACCAGA TGCGGTGTGA AATACCGCAC
10451	AGATGCGTAA GGAGAAAATA CCGCATCAGC CTCCATTCGC CATTCAGACT
10501	CCGCAACTGT TGGGAAGGGC GGTCAGTACG CGCTTCTTCG CTATTACGCC
10551	AACTGGCGAA AGGGGGATGT GCTGCAAGGC GATTAAGTTG GGTAACGCCA
10601	GGGTTTTCCC AATCACGACG TTGTAAAACG ACAGCCAATG AATTGAAGCT
10651	TATTAATTCT AGACTGAAGC TTTTAATACG ACTCACTATA (SEQ. ID. NO.:3)

Fig. 1G

#### SEQUENCE LISTING

<110> Istituto Di Ricerche Di Biologia Molecolare P. Angeletti S.P.A.
<120> HEPATITIS C VIRUS REPLICONS AND REPLICON
ENHANCED CELLS
<130> IT0003 PCT
<150> 60/263,479
<151> 2001-01-23

<170> FastSEQ for Windows Version 4.0

<160> 13

<210> 1
<211> 3010
<212> PRT
<213> Con 1 HCV isolate nucleic acid

<400> 1 Met Ser Thr Asn Pro Lys Pro Gln Arg Lys Thr Lys Arg Asn Thr Asn Arg Arg Pro Gln Asp Val Lys Phe Pro Gly Gly Gly Gln Ile Val Gly Gly Val Tyr Leu Leu Pro Arg Arg Gly Pro Arg Leu Gly Val Arg Ala Thr Arg Lys Thr Ser Glu Arg Ser Gln Pro Arg Gly Arg Arg Gln Pro Ile Pro Lys Ala Arg Gln Pro Glu Gly Arg Ala Trp Ala Gln Pro Gly Tyr Pro Trp Pro Leu Tyr Gly Asn Glu Gly Leu Gly Trp Ala Gly Trp Leu Leu Ser Pro Arg Gly Ser Arg Pro Ser Trp Gly Pro Thr Asp Pro Arg Arg Arg Ser Arg Asn Leu Gly Lys Val Ile Asp Thr Leu Thr Cys Gly Phe Ala Asp Leu Met Gly Tyr Ile Pro Leu Val Gly Ala Pro Leu Gly Gly Ala Ala Arg Ala Leu Ala His Gly Val Arg Val Leu Glu Asp Gly Val Asn Tyr Ala Thr Gly Asn Leu Pro Gly Cys Ser Phe Ser Ile Phe Leu Leu Ala Leu Leu Ser Cys Leu Thr Ile Pro Ala Ser Ala Tyr Glu Val Arg Asn Val Ser Gly Val Tyr His Val Thr Asn Asp Cys Ser Asn Ala Ser Ile Val Tyr Glu Ala Ala Asp Met Ile Met His Thr Pro Gly Cys Val Pro Cys Val Arg Glu Asn Asn Ser Ser Arg Cys Trp Val Ala Leu Thr Pro Thr Leu Ala Ala Arg Asn Ala Ser Val Pro Thr Thr Thr Ile Arg Arg His Val Asp Leu Leu Val Gly Ala Ala Ala Leu Cys Ser Ala Met Tyr Val Gly Asp Leu Cys Gly Ser Val Phe Leu Val Ala Gln Leu Phe Thr Phe Ser Pro Arg Arg His Glu Thr Val Gln Asp Cys . 295 Asn Cys Ser Ile Tyr Pro Gly His Val Thr Gly His Arg Met Ala Trp 

Asp Met Met Asn Trp Ser Pro Thr Ala Ala Leu Val Val Ser Gln Leu Leu Arg Ile Pro Gln Ala Val Val Asp Met Val Ala Gly Ala His Trp Gly Val Leu Ala Gly Leu Ala Tyr Tyr Ser Met Val Gly Asn Trp Ala Lys Val Leu Ile Val Met Leu Leu Phe Ala Gly Val Asp Gly Gly Thr Tyr Val Thr Gly Gly Thr Met Ala Lys Asn Thr Leu Gly Ile Thr Ser Leu Phe Ser Pro Gly Ser Ser Gln Lys Ile Gln Leu Val Asn Thr Asn Gly Ser Trp His Ile Asn Arg Thr Ala Leu Asn Cys Asn Asp Ser Leu Asn Thr Gly Phe Leu Ala Ala Leu Phe Tyr Val His Lys Phe Asn Ser Ser Gly Cys Pro Glu Arg Met Ala Ser Cys Ser Pro Ile Asp Ala Phe Ala Gln Gly Trp Gly Pro Ile Thr Tyr Asn Glu Ser His Ser Ser Asp Gln Arg Pro Tyr Cys Trp His Tyr Ala Pro Arg Pro Cys Gly Ile Val Pro Ala Ala Gln Val Cys Gly Pro Val Tyr Cys Phe Thr Pro Ser Pro Val Val Gly Thr Thr Asp Arg Phe Gly Val Pro Thr Tyr Ser Trp Gly Glu Asn Glu Thr Asp Val Leu Leu Leu Asn Asn Thr Arg Pro Pro Gln Gly Asn Trp Phe Gly Cys Thr Trp Met Asn Ser Thr Gly Phe Thr Lys Thr Cys Gly Gly Pro Pro Cys Asn Ile Gly Gly Ile Gly Asn Lys Thr Leu Thr Cys Pro Thr Asp Cys Phe Arg Lys His Pro Glu Ala Thr Tyr Thr Lys Cys Gly Ser Gly Pro Trp Leu Thr Pro Arg Cys Leu Val His Tyr Pro Tyr Arg Leu Trp His Tyr Pro Cys Thr Val Asn Phe Thr Ile Phe Lys Val Arg Met Tyr Val Gly Gly Val Glu His Arg Leu Glu Ala Ala Cys Asn Trp Thr Arg Gly Glu Arg Cys Asn Leu Glu Asp Arg Asp Arg Ser Glu Leu Ser Pro Leu Leu Leu Ser Thr Thr Glu Trp Gln Val Leu Pro Cys Ser Phe Thr Thr Leu Pro Ala Leu Ser Thr Gly Leu Ile His Leu His Gln Asn Val Val Asp Val Gln Tyr Leu Tyr Gly Ile Gly Ser Ala Val Val Ser Phe Ala Ile Lys Trp Glu Tyr Val Leu Leu Leu Phe Leu Leu Ala Asp Ala Arg Val Cys Ala Cys Leu Trp Met Met Leu Leu Ile Ala Gln Ala Glu Ala Ala Leu Glu Asn Leu Val Val Leu Asn Ala Ala Ser Val Ala Gly Ala His Gly Ile Leu Ser Phe Leu Val Phe Phe Cys Ala Ala Trp Tyr Ile Lys Gly Arg Leu Val Pro Gly Ala Ala Tyr Ala Leu Tyr Gly Val Trp Pro Leu Leu Leu Leu Leu Leu Ala Leu Pro Pro Arg Ala Tyr Ala Met Asp Arg Glu Met Ala Ala 

Ser Cys Gly Gly Ala Val Phe Val Gly Leu Ile Leu Leu Thr Leu Ser Pro His Tyr Lys Leu Phe Leu Ala Arg Leu Ile Trp Trp Leu Gln Tyr Phe Ile Thr Arg Ala Glu Ala His Leu Gln Val Trp Ile Pro Pro Leu Asn Val Arg Gly Gly Arg Asp Ala Val Ile Leu Leu Thr Cys Ala Ile His Pro Glu Leu Ile Phe Thr Ile Thr Lys Ile Leu Leu Ala Ile Leu Gly Pro Leu Met Val Leu Gln Ala Gly Ile Thr Lys Val Pro Tyr Phe Val Arg Ala His Gly Leu Ile Arg Ala Cys Met Leu Val Arg Lys Val Ala Gly Gly His Tyr Val Gln Met Ala Leu Met Lys Leu Ala Ala Leu .930 Thr Gly Thr Tyr Val Tyr Asp His Leu Thr Pro Leu Arg Asp Trp Ala His Ala Gly Leu Arg Asp Leu Ala Val Ala Val Glu Pro Val Val Phe Ser Asp Met Glu Thr Lys Val Ile Thr Trp Gly Ala Asp Thr Ala Ala 980 985 990 Cys Gly Asp Ile Ile Leu Gly Leu Pro Val Ser Ala Arg Arg Gly Arg Glu Ile His Leu Gly Pro Ala Asp Ser Leu Glu Gly Gln Gly Trp Arg Leu Leu Ala Pro Ile Thr Ala Tyr Ser Gln Gln Thr Arg Gly Leu Leu Gly Cys Ile Ile Thr Ser Leu Thr Gly Arg Asp Arg Asn Gln Val Glu Gly Glu Val Gln Val Val Ser Thr Ala Thr Gln Ser Phe Leu Ala Thr Cys Val Asn Gly Val Cys Trp Thr Val Tyr His Gly Ala Gly Ser Lys Thr Leu Ala Gly Pro Lys Gly Pro Ile Thr Gln Met Tyr Thr Asn Val 1090 1095 1100 Asp Gln Asp Leu Val Gly Trp Gln Ala Pro Pro Gly Ala Arg Ser Leu Thr Pro Cys Thr Cys Gly Ser Ser Asp Leu Tyr Leu Val Thr Arg His 1125 1130 1135 Ala Asp Val Ile Pro Val Arg Arg Gly Asp Ser Arg Gly Ser Leu Leu Ser Pro Arg Pro Val Ser Tyr Leu Lys Gly Ser Ser Gly Gly Pro Leu Leu Cys Pro Ser Gly His Ala Val Gly Ile Phe Arg Ala Ala Val . 1180 Cys Thr Arg Gly Val Ala Lys Ala Val Asp Phe Val Pro Val Glu Ser Met Glu Thr Thr Met Arg Ser Pro Val Phe Thr Asp Asn Ser Ser Pro . 1215 Pro Ala Val Pro Gln Thr Phe Gln Val Ala His Leu His Ala Pro Thr Gly Ser Gly Lys Ser Thr Lys Val Pro Ala Ala Tyr Ala Ala Gln Gly Tyr Lys Val Leu Val Leu Asn Pro Ser Val Ala Ala Thr Leu Gly Phe Gly Ala Tyr Met Ser Lys Ala His Gly Ile Asp Pro Asn Ile Arg Thr Gly Val Arg Thr Ile Thr Thr Gly Ala Pro Ile Thr Tyr Ser Thr Tyr Gly Lys Phe Leu Ala Asp Gly Gly Cys Ser Gly Gly Ala Tyr Asp Ile 

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